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(57) Abstract

There are provided according to the invention, novel compounds of formula (I) wherein R¹, R², R³, R⁴, R⁵ and R⁶ are as defined in the specification, processes for preparing them, formulations containing them and their use in therapy for the treatment of inflammatory diseases.

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Compounds useful in the treatment of inflammatory diseases

This invention relates to novel chemical compounds, processes for their preparation, pharmaceutical formulations containing them and their use in therapy.

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Inflammation is a primary response to tissue injury or microbial invasion and is characterised by feukocyte adhesion to the endothelium, diapedesis and activation within the tissue. Leukocyte activation can result in the generation of toxic oxygen species (such as and lymphocytes. Different forms of inflammation involve different types of infiltrating superoxide anion), and the release of granule products (such as peroxidases and Circulating leukocytes include neutrophils, eosinophils, basophils, monocytes leukocytes, the particular profile being regulated by the profile of adhesion molecule, cytokine and chemotactic factor expression within the tissue.

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The primary function of leukocytes is to defend the host from invading organisms, such as causes the local recruitment of leukocytes from the circulation into the affected tissue. However in chronic inflammatory states, recruitment is often inappropriate, resolution is not bacteria and parasites. Once a tissue is injured or infected, a series of events occurs which Leukocyte recruitment is controlled to allow for the orderly destruction and phagocytosis of foreign or dead cells, followed by tissue repair and resolution of the inflammatory infiltrate. adequately controlled and the inflammatory reaction causes tissue destruction.

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than neutrophils (eg. eosinophils, T- and B-lymphocytes, basophils and mast cells). The (CS-1) in fibronectin (an extracellular matrix protein), and (iii) a site on the mucosal minimally expressed in the vasculature, however, upregulation of VCAM-1 on endothelial. and some cortical neurons. MAdCAM expression is predominantly associated with gut tissue Integrins are cell surface heterodimeric proteins comprising α and β chains, involved in the inflammatory process. The $\alpha 4$ -integrins, which include $\alpha 4\beta 1$ (also known as very late antigen-4 (VLA-4) or CD49d/CD29) and $\alpha4\beta7$, are expressed mainly on leukocytes other (VCAM-1; CD106), (ii) a sequence within the alternatively spliced connecting segment-1 addressin cell adhesion molecule (MAdCAM). Under normal conditions, VCAM-1 is cells occurs near sites of Inflammation. VCAM-1 has also been identified on a range of nonvascular cells including dendritic cells, bone marrow stromal cells, synoviocytes, astrocytes adhesion molecule ligands for $\alpha 4$ -integrins include (i) the vascular cell adhesion molecule

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being expressed in the high endothelial veins of gut associated lymphold tissue, peripheral lymph nodes and Peyers Patches.

eukocytes through the wall of post-capillary venules to sites of tissue inflammation. Such an are capable of intervention at two levels to effect attenuation of the inflammatory processes nhibition of the recrultment of leukocytes to sites of tissue inflammation and (ii) inhibition of The $\alpha 4$ -integrin/VCAM-1 interaction enables adhesion and subsequent transmigration of interaction is similarly capable of providing a co-stimulatory signal for T-cell activation, whilst Both a4B1 (VLA-4) and a4B7 can interact with VCAM-1, CS-1 in fibronectin and MAdCAM. degranulation of mast cells, basophils and eosinophils. Therefore, a4-integrin antagonists roje which are essential in the pathophysiology of many chronic diseases. These a stimulatory the activation of leukocytes and the release of inflammatory mediators. the a4-integrin/fibronectin interaction is believed to have

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Cell adhesion and signalling, mediated by $\alpha 4$ -integrins, are essential in numerous (Lobb and Hemler, 1994). Anti-a4-mAbs have shown beneficial effects in animal models of allergic lung inflammation relevant to asthma, including guinea-pig, rat, rabbit and sheep models. Additionally, anti-α4-mAbs have also been shown to be efficacious in (i) rat and mouse models of experimental allergic encephalomyelitis (considered to be a model of the hypersensitivity, (iii) colitis in the Cotton-top tamarin, relevant to inflammatory bowel disease (Podolsky et al, 1993), and (iv) insulin dependent diabetes melitus in the non-obese diabetic physiological and pathophysiological processes. The therapeutic potential of α4-integrin blocking agents has been investigated previously by testing specific a4-integrin blocking I-cell dependent autoimmune disease, multiple scierosis), (ii) mouse models of contact monoclonal antibodies (anti-a4-mAbs) in experimental in vitro and in vivo models of disease mouse (Baron et al, 1994). Fibronectin-derived peptides which are thought to block lpha 4integrin function have shown efficacy in mouse contact hypersensitivity (Ferguson et al, 1991) and in rat adjuvant arthritis (Wahl et al, 1994). 15 20

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International patent application numbers WO 98/53814, WO 98/53817 and WO 98/53818 (Merck) describe the use of heterocyclic amide compounds, biaryfalkanoic aclds and sulphonamide compounds, respectively, as VLA-4 and/or $\alpha 4/\beta 7$ antagonists. WO 98/54207 (Celltech) describes the use of tyrosine derivatives to inhibit the binding of $\alpha4$ integrins to their ligands for the treatment and prophylaxis of immune or anti-inflammatory disorders.

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WO97/03094 (Blogen) describes a selection of semi-peptidic compounds which are capable of inhibiting the binding of ligands to the VLA-4 receptor.

We have now found a novel group of $\alpha 4$ -integrin antagonist compounds which antagonise both $\alpha 4\beta 1$ and $\alpha 4\beta 7$ integrins, with the potential to block feukocyte adhesion and activation, consequently effecting anti-inflammatory properties. These compounds are therefore of potential therapeutic benefit, especially in providing protection from leukocyte-induced tissue damage in diseases where leukocytes are implicated at the site of inflammation. Antagonists of both $\alpha 4\beta 1$ and $\alpha 4\beta 7$ integrins may have advantages over selective antagonists of $\alpha 4\beta 1$ or $\alpha 4\beta 7$ because both integrins are believed to have a role in inflammation.

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Thus, according to one aspect of the invention, we provide compounds of formula I:

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wherein R1 and R2 independently represent

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(i) -C₁₋₆ alkyl, -C₃₋₆ cycloalkyl or -C₁₋₃ alkylC₃₋₆ cycloalkyl, or such a group in which alkyl or cycloalkyl is substituted by one or more halogen, -CN, nitro, hydroxy or -OC₁₋₆alkyl groups;

(ii) -(CH2),Ar' or -(CH2),OAr';

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-(CH₂),COOR"; -(CH₂),Ar', -O(CH₂),Ar', -(CH₂),CO(CH₂),Ar' or -(CH₂),OAr';

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or R³ represents -(CH₂)₀-2,4-imidazolidinedione, -(CH₂)₀(piperidin-4-yl), -(CH₂)₀(piperidin-3-yl), -(CH₂)₀(piperidin-2-yl), -(CH₂)₀(piperidin-2-yl), -(CH₂)₀(piperidin-2-yl), -(CH₂)₀(piperidin-2-yl), -(CH₂)₀(morpholin-3-yl) or -(CH₂)₀(morpholin-2-yl) optionally substituted on nitrogen by -(CO)₀C₁₂alkyl, -(CO)₀(CH₂)₀Ar² or -C(=NH)NH₂;

or R³ represents -(CH₂)₂dibenzofuran optionally substituted by -C₁₂alkyl or halogen; or R³ represents -(CH₂)₂-thioxanthen-9-one;

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R* represents hydrogen, -C₁₋₈ alkyl, -C₁₋₈ alkylC₃₋₆ cycloalkyl, -(CH₂), Ar², -C₁₋₄alkyl-X-R², -C₁₋₄alkyl-X-R², -C₁₋₄alkyl SO₂C₁₋₄ alkyl, -C₁₋₄alkylNR¹²R¹³ or -C₁₋₄alkylNR¹²COC₁₋₆ alkyl; R⁵ represents hydrogen, or R⁴R⁵ together with the carbon to which they are attached form a C₆₋₇ cycloalkyl ring;

Re represents hydrogen or "C₁₋₈alkyl, or Re and R* together with the N and C atoms to which they are respectively attached form a pyrrolidine ring;

R' represents hydrogen, -(CH2),NR12R13, -(CH2),Ar2 or -(CH2),NR12COC, alkyl;

R°, R°, R¹6 and R¹7 independently represent hydrogen, -C₁-aalkyl, -C₃-cycloalkyl, -C₃-cycloalkyl, -C₁-a alkylC₃-6 cycloalkyl, -C₂-aalkenyl or NR6R® or NR¹6R¹ together represents morpholinyl, pyrrolidinyl, piperazinyl or piperazinyl N-substituted by -C₁-a alkyl, -COphenyl or

SO₂methyl;

R¹⁰, R¹¹, R¹², R¹³, R¹⁶, R²⁰ and R²¹ independently represent hydrogen or -C₁₋₆alkyl;

R¹⁰, R¹¹, R¹² and R²² independently represent hydrogen, -C₁₋₆alkyl, -C₃₋₆ cycloalkyl or -(CH₂)_x Ar⁴ or NR¹⁴R¹⁶ or NR¹⁵R²² together represents morpholinyl, pyrrolidinyl, piperidinyl, piperazinyl or N-C₁₋₆alkylpiperazinyl;

Ar¹ represents phenyl or a 5 or 8 membered heterocyclic aromatic ring containing 1 to 3 heteroatoms selected from O, N and S optionally substituted by one or more halogen, C₁₂alkyl; hydroxy, -OC₁₂alkyl, CF₃, nitro, -Ar² or -OAr² groups;

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Ar² represents phenyl optionally substituted by one or more halogen, -C_{1-s}alkyl, hydroxy, -OC_{1-s}alkyl, -CF₃ or nitro groups;

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Ar represents phenyl, a 5 or 6 membered heterocyclic aromatic ring containing 1 to 3 heteroatoms selected from O, N or S, or such a group fused to a benzene ring, optionally substituted by one or more -CO(CH₂)₀Ar', -(CH₂)_yAr', -(CH₂)_yCOAr', -(CO)_aC₁₋₆ alkyl, -(CO)_aC₂₋₆ alkenyl, -(CO)_aC₂₋₆ alkynyl, -(CO)_aC₃₋₆ cyctoalkyl, -(CO)_aC₁₋₆ haloalkyl, halogen,

-COCH₂CN, -(CH₂)_bNR¹⁶R¹⁷, -(CH₂)_bNHC(=NH)NH₂, -CYNR¹⁶(CO)_aR¹⁷, -(CH₂)_bNR¹⁵COR¹⁸, -(CH₂)_bCONR¹⁶R²², -(CH₂)_bNR¹⁵CONR¹⁶(CH₂)_bNR¹⁵CONR¹⁶, -SO₂R¹⁸, -SO₂R¹⁹, -SO₇R¹⁹, -(CH₂)_bOH, -COOR¹⁶, -CHO, -OC₁, roalkyl, -O(CH₂)_fNR¹⁵R²², -O(CH₂)_fNHC(=NH)NH₂, -O(CH₂)_bCONR¹⁶, -O(CH₂)_bCONR¹⁶, -O(CH₂)_bCONR¹⁶, -O(CH₂)_bCONR¹⁶, -O(CH₂)_bCONR¹⁶, -O(CH₂)_bOAr², -O(CH₂)_bAr², 3-phenyl-2-pyrazotin-5-one or 4,5-dihydro-3(2H)-pyrldazinone groups;

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Ar represents phenyl or a 5 or 6 membered heterocyclic aromatic ring containing 1 to 3 heteroatoms selected from O, N and S optionally substituted by one or more halogen, -C₁₋₆alkyl, hydroxy, -OC₁₋₆alkyl, -CF₃, nitro or -CONH₂ groups;

X and Y independently represent O or S;

a, f, k, s and n independently represent 0 or 1;

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b, c, r, x, y and z independently represent an integer 0 to 2;

d, g and u independently represent 1 or 2;

e, h, q and w independently represent an integer 1 to 3;

Jand p Independently represent an integer 2 to 4;

m independently represents an integer 0 to 4;

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t independently represents an integer 0 to 3;

and salts and solvates thereof.

Examples of 5 or 6 membered heterocyclic aromatic rings that Ar¹, Ar³ and Ar⁴ may represent include pyrlmldine, pyridine, furan, imidazole, thiophene, pyrrole, thiazole, oxazole isoxazole, 1,3,4-thiadiazole, 1,2,3-thiadiazole, 1,2,4-thiadiazole, 1,2,4,4-thiadiazole, 1,2,4,4-thiadiazole, 1,2,4,4-th

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Specific examples of 5 or 6 membered heterocyclic aromatic rings that Ar¹ may represent include pyrimidine, pyridine, furan, 1,2,4-thiadiazole and pyrrole.

Specific examples of 5 or 6 membered heterocyclic aromatic rings that Ar³ may represent include thiazole and pyridine. Phenyl fused to a benzene ring represents naphthyl. An example of a 5 or 6 membered heterocyclic aromatic ring fused to a benzene ring that Ar³ may represent includes benzofuran.

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Specific examples of 5 or 6 membered heterocyclic aromatic rings that Arf may represent lnclude 1,3,4-thiadiazole, 1,2,3-thiadiazole, 1,2,4-oxadiazole and pyrazole.

We prefer R¹ and R² to be defined such that NR¹R² together represent piperidinyl, piperazinyl, thiomorpholinyl, morpholinyl or 1,2,3,4-tetrahydrolsoquinoline optionally substituted by a -(CO), (CH₂),Ar¹, -(CO),C₁,alkyl, -(CH₂),CONR⁴R², -NR¹⁰(CO),(CH₂),Ar¹, -NR¹⁰(CO),C₁,a alkylC₃,a cycloalkyl, -(CH₂),OC₁,a alkyldiC₃,a cycloalkyl, -(CH₂),OC₁,a alkyl, -(CH₂),O(CH₂),OH, piperidin-1-yl, -(CH₂),OH or -CONR⁴⁰(CH₂),Ar¹ group.

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We particularly prefer R¹ and R² to be defined such that NR¹R² together represents morpholinyl or piperazinyl optionally N-substituted by -(CO),C₁, alkyl (especially -COCH₃), piperazinyl N-substituted by -(CO),(CH₂),Ar¹ (especially -COphenyl and -(CO)₂-furanyl),

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piperidinyl substituted by -NR¹º(CO)"(CH₂),Ar¹ (especially -NHCOCH₂phenyl) or piperidinyl substituted by -(CH₂),CONRªR³ (especially -CONH₂).

We prefer R³ to represent -(CH₂),-2,4-imidazolidinedione-3-yt, -(CH₂),-thioxanthen-9-one-3-yt, -(CH₂),Ar³, -O(CH₂),Ar³, -(CH₂),OAr³ or -(CH₂),dibenzofuran, particularly -OCH₂Ar², -CH₂OAr³ or dibenzofuran, especially -CH₂OAr³ or dibenzofuran.

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When R³ represents -{CH₂},dibenzofuran (particularly dibenzofuran), we prefer it to represent -{CH₂},-2-dibenzofuran (particularly 2-dibenzofuran).

When R³ represents -(CH₂)_c-2,4-imidazolidinedione, we prefer it to represent -(CH₂)_c-(2,4-imidazolidinedione-3-yl) (particularly -CH₂-2,4-imidazolidinedione-3-yl).

When R³ represents -(CH₂)c-thioxanthen-9-one, we prefer it to represent -(CH₂)c-(thioxanthen-9-one-3-yl) (particularly -CH₂-thioxanthen-9-one-3-yl). We most especially prefer R³ to represent -CH₂OAr³.

We prefer R⁴ to represent -C₁₋₈ alkyl, R⁵ to represent hydrogen or for R⁴R⁵, together with the carbon to which they are attached, to form a cyclohexyl ring, and for R⁶ to represent hydrogen or methyl (particularly hydrogen).

We particularly prefer R⁴ to represent -C₁₋₆ alkyl, and for R⁵ and R⁶ to represent hydrogen. We especially prefer R⁴ to represent -CH₂CHMe₂ and for R⁵ and R⁶ to represent hydrogen. We particularly prefer R⁴ and R⁵ to have the stereochemical orientation shown in formula (la):

We prefer R7 to represent -(CH2),Ar2 or -(CH2),NR12COC1.6 alkyl.

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We especially prefer R⁸ and R⁹ each to represent hydrogen or for NR⁹R⁹ together to represent piperidinyl or pyrrolidinyl, particularly piperidinyl.

We prefer R¹⁰ to represent hydrogen or methyl, particularly hydrogen.

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We prefer R¹¹ to represent hydrogen or methyl, particularly hydrogen.

We prefer R¹² to represent hydrogen or methyl, particularly hydrogen.

We prefer R¹³ to represent hydrogen or methyl, particularly hydrogen.

We prefer R¹⁴ to represent hydrogen or methyl, particularly hydrogen.

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We prefer R¹⁶ to represent hydrogen, -C₁₄ alkyl or -C₂₄ alkenyl, particularly hydrogen or

We prefer R^{17} to represent hydrogen, - C_{14} alkyl or - C_{24} alkenyl, particularly hydrogen, meth)

We prefer R18 to represent hydrogen or methyl, particularly hydrogen. S

We prefer R¹⁹ to represent hydrogen or -C₁₋₈ alkyl, particularly -C₁₋₈ alkyl, especially methyl.

We prefer \mathbb{R}^2 to represent hydrogen or methyl, particularly hydrogen.

We prefer R^{24} to represent hydrogen or methyl, particularly hydrogen.

We prefer R22 to represent hydrogen, -C14 alkyl or -(CH2),Ar4 or for NR16R22 together to

represent piperidinyl, pyrrolidinyl or morpholinyl. 10

We especially prefer R15 and R22 to be defined such that NR15R22 together represents

We prefer Art to represent furan, pyrimidine or phenyl optionally substituted by halogen (eg. chlorine or fluorine) or -OC1-6 alkyl.

We prefer Art to represent unsubstituted phenyl. 15

We prefer Ar to represent phenyl, naphthyl or benzofuran optionally substituted by one or more -(CH2),COAr', -COOR15, -(CH2),SO2NR15R22, -(CH2),NR15SO2R19, -SO2R19, (CO),C24 pyrazolin-5-one-2-yi or 4,5-dihydro-3(2H)-pyrldazinone-6-yi groups. We particularly prefer alkenyl, -(CO), C, alkyl, -(CO), C, cycloalkyl, halogen, -(CH2), CONR 18R2, 3-phenyl-2-

Ar to represent phenyl or naphthyl optionally substituted by -(CO) C1-8 alkyl, -(CO) C3-8 cycloalkyl, halogen, -(CH2),COAr or -(CH2),CONR16R22. 20

cyclohexyl, todine, -COphenyl or COpiperidin-1-yl or naphthyl substituted by COpiperidin-1-We most particularly prefer Ar³ to represent phenyl substituted by n-propyl, tertiary butyl,

We prefer Art to represent phenyl or furan optionally substituted by halogen, especially unsubstituted phenyl or furan. 25

We prefer e to represent 1 or 2.

We prefer n to represent 0 or 1.

We prefer r to represent 0 or 1, particularly 1.

We prefer p to represent 2. 30

We prefer t to represent 0, 1 or 3, particularly 0 or 1, especially 0.

We prefer h to represent 1 or 2, particularly 2.

We prefer d to represent 1.

We prefer m to represent 0 or 1, particularly 1.

We prefer c to represent 0 or 1, particularly 1.

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We prefer f to represent 1.

We prefer q to represent 1 or 2, particularly

S

We prefer b to represent 0 or 1, particularly 0.

We prefer J to represent 2 or 3, particularly 2.

We prefer z to represent 0 or 1, particularly 0. 10

We prefer k to represent 1.

The most preferred compounds of formula (I) are:

(2S)-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methyl pentanoyl)amino}-3-(4-[(4morpholinylcarbonyl)oxylphenyl)propanoic acid; (2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-{4-[(4morpholinylcarbonyl)oxy]phenyl}propanoic acid; 20

(2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{[(2S)-2-((2-{2-(tertbutyl)phenoxy]acetyl}amino)-4-methyipentanoyl]amino)propanoic acid; (2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-{(4morpholinylcarbonyl)oxy]phenyl}propanoic acid;

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2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino}-3-{4-[(4morpholinylcarbonyl)oxy]phenyl}propanoic acid;

(2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino)-3-{4-{({4-((2-phenylacetyl)amino]-1-piperidinyl)carbonyl) oxy] phenyl)propanoic acid;

[2S]-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-[((2S)-4-methyl-2-{[2-(2methylphenoxy)acetyljamino}pentanoyl)amino]propanoic acid;

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(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{[(2S)-2-{(2-{2-(tert-

butyl)phenoxy]acetyl}amino)-4-methylpentanoyl]amino)propanoic acid;

(2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-({(2S)-2-[(dibenzo[b,d)furan-4-

/Icarbonyl)amino]-4-methytpentanoyl)amino)propanoic acid; 35

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We prefer u to represent 1.

We prefer w to represent 1 or 2, particularly 1.

We prefer x to represent 0 or 1, particularly 1.

We prefer a to represent 0.

We prefer y to represent 0 or 1, particularly 0.

We prefer s to represent 0.

We prefer g to represent 1.

We prefer X to represent oxygen.

We prefer Y to represent oxygen. 15

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(2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-[4-({[4- (2-furoyi)-1-plperazinyl]carbonyl}oxy)phenyl] propanoic acid;

(2S)-2-(((2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl}amino)-3-[4-({[4-((2s)-2-((2s)-1-piperazinyl]carbonyl}oxy)phenyl] propanoic acid;

(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{((2S)-4-methyl-2-{[2-(2-methylphenoxy)acetyl]amino}pentanoyl)amino]propanoic acid;

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(2S)-3-[4-([[4-(Aminocarbonyl]-1-piperidinyl]carbonyl]oxy)phenyl]-2-[((2S)-4-methyl-2-([2-(2-methylphenoxy)acetyl]amino)pentanoyl)amino] propanoic acid;

(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl}-2-{{(2S)-2-{(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl}amino)propanolc acld;

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(2S)-3-[4-(([4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-(((2S)-2-(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl}amino) propanoic acid; and salts and solvates thereof.

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(2S)-3-[4-([[4-(Aminocarbonyl)-1-plperidinyl]carbonyl)oxy)phenyl]-2-[[(2S)-2-({2-[2-(tert-butyl)phenoxy]acetyl)amino)-4-methylpentanoyl]amino} propanoic acid; and salts and solvates thereof.

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The above preferred compounds are characterised by low oral bioavailability which is an advantageous property for an inhaled medicine in order to minimise potential side effects.

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Suitable salts of the compounds of formula (I) include physiologically acceptable salts such as alkali metal salts, for example calcium, sodium and potassium salts and salts with (trishydroxymethyl)aminomethane. Other salts of the compounds of formula (I) include salts which may not be physiologically acceptable but may be useful in the preparation of compounds of formula (I) and physiologically acceptable salts thereof. If appropriate, acid addition salts may be derived from inorganic or organic acids, for example hydrochlorides, hydrobromides, sulphates, phosphates, acetates, benzoates, citrates, succinates, lactates, tartrates, furnarates, maleates, 1-hydroxynathanoate, methanesulphonate. Examples of solvates include hydrates.

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When sidechains of compounds of formula (I) contain chiral centres, the invention extends to mixtures of enantiomers (including racemic mixtures) and diastereoisomers as well as to Individual enantiomers. Generally it is preferred to use a compound of formula (I) in the form

of a purified single enantiomer.

The compounds of formula (I) and salts and solvates thereof may be prepared by the methodology described hereinafter, constituting a further aspect of this invention.

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A process according to the Invention for preparing a compound of formula (I) comprises;

(a) hydrolysis of a carboxylic acid ester of formula (ii)

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wherein R¹, R², R³, R⁴ and R⁵ are as defined above and R is a group capable of forming a carboxylic acid ester, or

deprotecting a compound of formula,(I) which is protected.

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In process (a) an example of a suitable R group is a C₁₄ alkyl group such as methyl or t-butyl. Hydrolysis may either occur via an acidic process e.g. involving trifluoroacetic acid and water or via.an alkaline route e.g. utilising sodlum hydroxide and methanol.

In an alternative solid phase reaction, R may represent a solid support functionalised with available hydroxy groups. Examples of solid supports include resins such as polystyrene resins wherein phenyl rings are provided with hydroxy groups via linkers. An example of a hydroxy functionalised linker is -CH₂O(4-hydroxymethyl-phenyl) (Wang Resin) or an N-Fmoc amino acid acyl ester of 3-methoxy-4-oxymethyl-phenoxymethylated 1% divinylbenzene cross-linked polystyrene (Sasrin resin).

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In process (b) examples of protecting groups and the means for their removal can be found in T. W. Greene 'Protective Groups in Organic Synthesis' (J. Wiley and Sons, 1991). Suitable amine protecting groups Include sulphonyl (e.g. tosyl), acyl (e.g. benzyloxycarbonyl or t-butoxycarbonyl) and arylalkyl (e.g. benzyl), which may be removed by hydrolysis or hydrogenolysis as appropriate.

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Compounds of formula (il) may be prepared following Scheme 1;

Step (i) In this Scheme we prefer R to represent methyl.

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Compounds of formula (III) and (IV) may be reacted under conventional conditions for preparation of an amide. Desirably a coupling agent eg. WSCDI with or without HOBT in an inert solvent such as MeCN or DMF is used. P, is an amine protecting group such as one described previously under process (b). In this Scheme we prefer P, to represent Boc.

Step (ii) The conversion of formula (V) to (VI) is suitably carried out with p-nitrophenyichloroformate under conventional conditions eg. in the presence of an organic base, eg. pyridine and an inert organic solvent such as DCM.

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Step (iii) This reaction may be performed by combination of the reagents in a suitable solvent, such as DCM in the presence of an organic base such as DIPEA.

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Step (Iv) This deprotection step may be performed under conventional conditions. When Prepresents Boc, it may be removed by treatment with acid e.g. a hydrohalic acid (HX) such as HCI:

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Step (v) A condensation reaction of formula (VIII) with the compound of formula R³CO₂H may be performed under conditions similar to those described above for step (i).

An alternative process for preparation of compounds of formula (II) is given in Scheme 2 below:

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Scheme 2

Step (i) In this Scheme we prefer R to represent t-Bu. The reaction conditions for this step are analogous to those for Scheme 1 step (i).

In compounds of formula (IV) in this Scheme we prefer P1 to represent Cbz.

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- Step (ii) This process comprises a two stage reaction, consisting of (a) treatment with a carboxyl donor such as (Cl₃CO)₂CO typically in the presence of an organic base such as DIPEA and a suitable solvent, such as THF or DCM followed by (b) conversion to the carbamate by treatment with R¹R²NH in a process analogous to that described previously in Scheme 1 step (iii).
- Step (iii) This deprotection reaction can be performed under conventional conditions. When P₁ represents Cbz, deprotection may be achieved by hydrogenolysis e.g. by treatment with ammonium formate in the presence of Pd/C in a solvent such as ethanol. The reaction may be worked up with acid, such as a hydrohalic acid to give the product as a hydrohalic acid salt (e.g. the HCl salt).

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Step (iv) This process is analogous to Scheme 1, step (v).

An alternative process for preparation of compounds of formula (II) is given in Scheme 3 below:

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Step (I) P₂ is an amine protecting group such as one described previously and in this Scheme we prefer P₂ to represent Fmoc. More preferably P₂ will be Boc.

A compound of formula (IX) may be reacted onto a suitable solid phase, such as a hydroxy functionalised polystyrene resin (e.g. Wang or Sasrin resin) in the presence of 2,6-dichlorobenzoyl chloride, pyridine and a suitable solvent, such as DMF.

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Step (ii) Removal of N-protecting group P₂ may be achieved under conventional conditions; e.g. when P₂ represents Fmoc, by treatment with an organic base such as piperidine in a suitable solvent, such as DMF or eg. when P₂ represents Boc, by treatment with chlorotrimethylsilane and phenol in a suitable solvent such as DCM.

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Step (iii) In this Scheme, P₁ may suitably represent Fmoc. Alternatively, it may suitably represent Boc. Reaction of a compound of formula (XI) with the compound of formula (IV) to produce an amide, may be performed in the presence of a coupling agent, such as PyBop, an organic base, such as DIPEA and a suitable solvent, such as DMF.

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Step (iv) This de-protection reaction may be performed under conventional conditions eg. when P₁ represents Finoc or Boc, under conditions analogous to those described above for

Step (v) A condensation reaction of formula (XIII) with the compound of formula R³CO₂H may be performed in the presence of a sultable coupling agent, such as PyBop, an organic base, such as DIPEA and a sultable solvent, such as DMF.

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Step (vI) This step comprises an alkenyl chain cleavage reaction on the compound of formula (XIV) to produce a compound of formula (XV), eg. by the treatment with Pd(PH₃), and PhSiH₃ (or morpholine) in the presence of a suitable solvent, such as DCM.

Step (vii) The conversion of a compound of formula (XV) to a compound of formula (XVI) is suitably performed by treatment with p-nitrophenyl chloroformate, under conventional conditions, in the presence of an organic base, such as DIPEA and an inert organic solvent, such as THF and/or DCM.

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Step (viii) This reaction may be performed by combination of the reagents in the presence of an organic base, such as DIPEA and suitable solvents, such as DCM and/or THF.

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An alternative process for preparation of certain compounds of formula (II) is given in Scheme 4 below:

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Step (!) In this Scheme we prefer P2 to represent Fmoc.

This conversion may be achieved following processes analogous to those of Scheme 3 steps (i) to (ii).

An alkenyl chain cleavage reaction may be performed by a process analogous to Scheme 3 step (vl). Step (ii)

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A p-nitrophenyl carbonate formation reaction, may be performed with reaction conditions analogous to Scheme 3 step (vil). Step (iii)

reaction Step (iv) The conversion of formula (XVIII) to (XIX) can be performed by analogous to Scheme 3 step (viii). This de-protection reaction may be performed using an analogous process to Scheme 3 step (II).

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Step (vi) The conversion of formula (XX) to (II) can be performed by a condensation reaction in the presence of a sultable acid, employing a sultable coupling agent, such as PyBop, an organic base, such as DIPEA and a solvent, such as DMF. Compounds of formula (II) in which R3 represents -(CH2)4OAr3 may alternatively be prepared from compounds of formula (XX) following steps (vil) and (viii): 9

Step (vil) The conversion of formula (XX) to (XXI) can be performed by a condensation reaction in the presence of a haloalkanoic acid (such as the bromo derivative i.e. Hal represents bromine), employing a suitable coupling agent, such as DIC and a solvent, such

Step (viii) In this step, the reaction of a compound of formula (XXI) with a compound of formula Ar-OH group may be undertaken in the presence of potassium carbonate, sodium iodide and a suitable solvent, such as DMF. Compounds of formula III, IV, IV, HNR'R2, R3COOH, IX, Hal(CH2)2COOH and AP-OH are either known or may be prepared by known methods.

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Compounds of the Invention may be tested for in vitro and in vivo biological activity in accordance with the following assays.

(1) Jurkat J6/VCAM-1 Adhesion Assay

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This assay was used to investigate the Interaction of the integrin VLA-4, expressed on the microtitre plates were coated with human immunoglobulin G (19G; Sigma Chemicals, UK, Polystyrene 96-well Jurkat J6 (human lymphoblast cell line) cell membrane with VCAM-1.

Product No. 14506) at a concentration of 0.05mg ml⁻¹ in bicarbonate buffer (36mM NaHCO₃ and 22mM Na₂CO₃, prepared in Dulbecco's phosphate buffered saline at pH 9.8 (PBS); Sigma Chemicals, UK, Product No. 14190-094) for 2 hours at 37°C. This solution was then aspirated and the plates were washed twice with PBS.

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vCAM-1 was prepared by cloning its constituent seven domains into a Drosophija expression system with a zz (Protein A) tag. This zzVCAM-1 was then expressed from Drosophija melanogaster S2 cell culture, induced with copper. Protease inhibitors were added and the culture supernatant was clarified either by filtration through a 0.2µm filter or by centrifugation. The zzVCAM-1 was then purified from this clarified medium using an IgG agarose column, equilibrated with either 20mM sodium phosphate pH 7.2 alone or in the presence of 0.5M sodium chloride. Elution of zzVCAM-1 from the column was mediated using 3M ammonium thiocyanate, which was subsequently removed using a G25 desalting column, equilibrated with 20mM sodium phosphate, pH 7.2. The purified zzVCAM-1 was then concentrated to a small volume (Amicon stirred cell concentrators) until a concentration of 62.5ng ml¹ was obtained, calculated using the extinction coefficient value.

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This solution of zzVCAM-1 was then incubated overnight at 4°C in the IgG coated microtitre plates with 3% bovine serum albumin (BSA) in PBS, followed by aspiration and two further washes with PBS. A concentration of the Jurkat J6 cells (6 x 10⁶ cells ml¹¹), grown in cell media RPMI 1640 (HyClone Ltd, Product No. B-9106-L) supplemented with 10% heat inactivated foetal calf serum (FCS; Gibco BRL, Product No. 10099-075) and 2mM L-glutamine, were labelled with 10µM of the fluorescent dye, 2', 7'-bis(2-carboxyethyl)-5-(e6)-carboxyfluoresceln acetoxymethyl ester (BCECF-AM; Molecular Probes Inc, Product No. B-1150) at 37°C for 10 minutes. The excess dye was then removed by centrifugation at 500xg for 5 minutes and the cells were resuspended at a concentration of 1.2·x 10⁷ cells mf¹ in Hank's balanced salt solution (HBSS; Gibco BRL, Product No. 14180-094).

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Equal volumes of compounds (dissolved in an appropriate solvent and diluted in HBSS containing 1mM MnCl₂) and the labelled Jurkat J6 cells, were added to the VCAM-1 coated plates and adhesion was allowed to proceed for 30 minutes at 37°C. Non, or loosely adhered cells, were removed by inversion of the plate and blotted with tissue paper. Two washes with PBS and further blotting were then performed, before the addition of 2% detergent (Triton-X®; Sigma Chemicals UK, Product No. X100). Counting was undertaken in a Wallac Viktor¹⁴ Fluorimeter, where low fluorescence values were indicative of compounds which had inhibited adhesion. All samples were assayed in singlicate and the following four parameter curve fit, shown by Equation (I) was applied:

Equation (I)

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Where a is the minimum, b is the Hill slope, c is the iC₅₀ and d is the maximum. (Maximum and minimum values are those compared to adhesion in the absence of compound and in the presence of the dipotassium salt of 2mM EDTA; Sigma Chemicals, UK, Product No. ED2P). Data is presented as the mean pIC₅₀ with the standard error of the mean of n experiments.

(2) CD3/VCAM-1 Co-stimulation of T-Lymphocyte Proliferation

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CD4* T-cells were purified from perlpheral blood mononuclear cells by negative selection with anti-CD14, CD19, CD16 and HLA.DR antibodies and Dynal beads. Flat bottomed 96-well tissue culture plates were coated with 1μg ml² anti-CD3 antibody (OKT3), washed and incubated with human IgG and zzVCAM-1 fusion proteins. The CD4* T cells (prepared in RPMI-1640 medium supplemented with 10% FCS, penicillin or streptomycln and L-glutamine) were added to the coated plates (1 × 10⁵ cells well*) and incubated in the presence or absence of various doses of compound or blocking antibodies for 4 days. Radiolabelled thymidine [³H] was added for the final 6 hours of incubation and the cells were then harvested using a Skatron plate harvester. Incorporation of the [³H] label was measured as an indicator of T cell proliferation using a β plate counter. Compounds were assayed in triplicate and data was collected in an analogous procedure to that described for Assay (1).

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(30 minutes breathing of an aerosol of the compound) or the intra-tracheal route, 30 minutes from a 0.5% solution of ovalbumin). Hyper-reactivity of the airways to the thromboxane sacrificed and the lungs lavaged. Total and differential leukocyte counts were then obtained before and 6 hours after ovalbumin challenge (10 minutes breathing of an aerosol generated animals using a whole body plethysmograph (Buxco Ltd., USA). The guinea pigs were then mimetic U46619, was measured 24 hours after ovalbumin challenge in un-restrained pigs were dosed with mepyramine (30mg kg.1 ip) to protect against anaphylactic bronchospasm. Test compounds, dissolved in 0.9% saline, were given by the inhaled route or the bronchoalveolar lavage fluid and the percentage reduction in eosinophil accumulation in a method based on that described by Danahay et el., 1997, ovalbumin sensitised gulnea control. Data was presented as the inhibitory effect of the specified dose expressed as determined (Sanjar et al., 1992). Dexamethasone (200µg kgr¹ i.t) was used as (3) Inhibition of Eosinophil Infiltration and Hyper-Reactivity in the Guinea Pig 20 25 30

(4) RPMI 8866/MAdCAM-1 Adhesion Assay

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percentage of the vehicle control response.

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This assay was used to investigate the interaction of the integrin α₄β₇, expressed on the RPMI 8866 (human B lymphold cell line) cell membrane with MAdCAM-1. Polystyrene 96-well microtitre plates were coated with human immunoglobulin G (IgG; Sigma Chemicals, UK, Product No. 14506) at a concentration of 0.05mg ml⁻¹ in bicarbonate buffer (36mM NaHCO₃ and 22mM Na₂CO₃, prepared in Dulbecco's phosphate buffered saline at pH 9.8 (PBS); Sigma Chemicals, UK, Product No. 14190-094) for 2 hours at 37°C. This solution was then aspirated and the plates were washed twice with PBS.

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MAdCAM-1 was prepared by cloning its constituent domains, under the control of a polyhedrin promoter, into a baculovirus expression system with a zz (Protein A) tag. The amplified baculovirus containing zzMAdCAM-1 was used to infect Spodoptera frugiperda cells growing in suspension in SF900II medium supplemented with 5% foetal calf serum. The cells were infected at a multiplicity of infection of 1 and harvested 48 hours later by centrifugation. Protease inhibitors were added and the culture supernatant was clarified either by filtration through a 0.2µm filter or by centrifugation. The zzMAdCAM-1 was then purified from this clarified medium using an IgG agarose column, equilibrated with either 20mM sodium phosphate pH 7.2 alone or in the presence of 0.5M sodium chloride. Elution of zzMAdCAM-1 from the column was mediated using 3M ammonium thiocyanate. The sample was then dialysed thoroughly, using 20mM sodium phosphate pH 7.2, to remove the ammonium thiocyanate. The purified zzMAdCAM-1 was then concentrated to a small volume (Amicon stirred cell concentrators) until a concentration of 0.5mg mf¹ was obtained, calculated using the extinction coefficient value.

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This solution of zzMAdCAM-1 was diluted 1:2500 and then incubated overnight at 4°C in the IgG coated microtitre plates with 3% bovine serum albumin (BSA) in PBS, followed by aspiration and two further washes with PBS. A concentration of the RPMI 8866 cells (3 x 10° cells ml¹¹), grown in cell media RPMI 1640 (HyClone Ltd, Product No. B-9106-L) supplemented with 10% heat inactivated foetal calf serum (FCS; Gibco BRL, Product No. 10099-075) and 2mM L-glutamine, were labelled with 10µM of the fluorescent dye, 2', 7'-bis(2-carboxyethyl)-5-(e6)-carboxyfluorescein acetoxymethyl ester (BCECF-AM; Molecular Probes Inc, Product No. B-1150) at 37°C for 10 minutes. The excess dye was then removed by centrifugation at 500xg for 5 minutes and the cells were resuspended at a concentration of 6 x 10° cells ml¹¹ in Hank's balanced salt solution (HBSS; Gibco BRL, Product No. 14190-

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Equal volumes of compounds (dissolved in an appropriate solvent and diluted in HBSS containing 1mM MnCl₂) and the labelled RPMI 8866 cells, were added to the MAdCAM-1 coated plates and adhesion was allowed to proceed for 30 minutes at 37°C. Non, or loosely adhered cells, were removed by inversion of the plate and blotted with tissue paper. Two washes with PBS and further blotting were then performed, before the addition of 2% detergent (Triton-X®; Sigma Chemicals UK, Product No. X100). Counting was undertaken in a Wallac Victor* Fluorimeter, where low fluorescence values were indicative of compounds which had inhibited adhesion. All samples were assayed in singlicate and the following four parameter curve fit, shown by Equation (I) (above) was applied. Wherein the maximum and minimum values are those compared to adhesion in the absence of compound and in the presence of the dipotassium salt of 2mM EDTA; Sigma Chemicals, UK, Product No. ED2P). Data is presented as the mean plC₆₀ with the standard error of the mean of n experiments.

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Examples of disease states in which the compounds of the invention have potentially beneficial anti-inflammatory effects include diseases of the respiratory tract such as bronchitis (including chronic bronchitis), asthma (including allergen-induced asthmatic reactions), chronic obstructive pulmonary disease (COPD) and rhinitis. Other relevant disease states include diseases of the gastrointestinal tract such as intestinal inflammatory diseases of the gastrointestinal tract such as intestinal inflammatory alseases secondary to radiation exposure or allergen exposure. Furthermore, compounds of the invention may be used to treat nephritis, skin diseases such as psoriasis, allergic dermatitis and hypersensitivity reactions and diseases of the central nervous system which have an inflammatory component eg. Alzheimer's disease, meningitis, multiple sclerosis and AIDS dementia.

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Further examples of disease states in which compounds of the invention have potentially beneficial effects include cardiovascular conditions such as atherosclerosis, peripheral vascular disease and idiopathic hypereoslnophilic syndrome.

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Compounds of the invention may be useful as immunosuppressive agents and so have use in the treatment of auto-immune diseases such as allograft tissue rejection after transplantation, rheumatold arthritis and diabetes.

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Compounds of the invention may also be useful in inhibiting metastasis.

Diseases of principal interest include asthma, COPD and inflammatory diseases of the upper respiratory tract involving seasonal and perennlal rhinitis.

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It will be appreciated by those skilled in the art that reference herein to treatment extends to prophylaxis as well as the treatment of established conditions.

As mentioned above, compounds of formula (I) are useful as pharmaceuticals, In particular as anti-inflammatory agents.

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There is thus provided as a further aspect of the invention a compound of formula (I) or physiologically acceptable saft or solvate thereof for use as pharmaceuticals, particularly the treatment of patients with inflammatory conditions.

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According to another aspect of the invention, there is provided the use of a compound of formula (I) or a physiologically acceptable salt or solvate thereof for the manufacture of a medicament for the treatment of patients with inflammatory conditions.

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In a further or alternative aspect there is provided a method for the treatment of a human or animal subject with an inflammatory condition, which method comprises administering to said human or animal subject an effective amount of a compound of formula (I) or a physiologically acceptable salt or solvate thereof.

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The compounds according to the invention may be formulated for administration in any convenient way, and the invention therefore also includes within its scope pharmaceutical compositions for use in anti-inflammatory therapy, comprising a compound of formula (i) or a physiologically acceptable saft or solvate thereof together, if desirable, with one or more physiologically acceptable diluents or carriers.

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There is also provided a process for preparing such a pharmaceutical formulation which comprises mixing the ingredients.

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The compounds according to the invention may, for example, be formulated for oral, buccat, parenteral, topical or rectal administration, preferably for topical administration to the lung, eg. by aerosol or as a dry powder composition.

Tablets and capsules for oral administration may contain conventional excipients such as binding agents, for example syrup, acacia, gelatin, sorbitol, tragacanth, mucilage of starch, cellulose or polyvinyl pyrrolidone; fillers, for example, factose, microcrystalline cellulose, sugar, maize- starch, calcium phosphate or sorbitol; lubricants, for example, magnesium stearate, stearic acid, talc, polyethylene glycol or silica; disintegrants, for example, potato starch, croscarmellose sodium or sodium starch glycollate; or wetting agents such as sodium

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lauryl sulphate. The tablets may be coated according to methods well known in the art. Oral liquid preparations may be in the form of, for example, aqueous or olly suspensions, solutions, emulsions, syrups or elixirs, or may be presented as a dry product for constitution with water or other sultable vehicle before use. Such liquid preparations may contain conventional additives such as suspending agents, for example, sorbitol syrup, methyl cellulose, glucose/sugar syrup, gelatin, hydroxymethyl cellulose, carboxymethyl cellulose, aluminium stearate gel or hydrogenated edible fats; emulsifying agents, for example, tecithin, sorbitan mono-oleate or acacia; non-aqueous vehicles (which may include edible oils), for example almond oil, fractionated coconut oil, oily esters, propylene glycol or ethyl alcohol; or preservatives, for example, methyl or propyl p- hydroxybenzoates or sorbic acid. The preparations may also contain buffer salts, flavouring, colouring and/or sweetening agents (e.g. mannitol) as appropriate.

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For buccal administration the compositions may take the form of tablets or lozenges formulated in conventional manner.

The compounds may also be formulated as suppositories, e.g. containing conventional suppository bases such as cocoa butter or other giycerides.

The compounds according to the invention may also be formulated for parenteral administration by bolus injection or continuous infusion and may be presented in unit dose form, for instance as ampoules, vials, small volume infusions or pre-filled syringes, or in multi-dose containers with an added preservative. The compositions may take such forms as solutions, suspensions, or emulsions in aqueous or non-aqueous vehicles, and may contain formulatory agents such as anti-oxidants, buffers, antimicrobial agents and/or tonicity adjusting agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g. sterile, pyrogen-free water, before use. The dry solid presentation may be prepared by filling a sterile powder aseptically into individual sterile containers or by filling a sterile solution aseptically into each container and freeze-drying.

By topical administration as used herein, we include administration by insufflation and inhalation. Examples of various types of preparation for topical administration include ointments, creams, totions, powders, pessaries, sprays, aerosols, capsules or cartridges for use in an inhaler or insufflator, solutions for nebulisation or drops (e.g. eye or nose drops).

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Ointments and creams may, for example, be formulated with an aqueous or oily base with the addition of suitable thickening and/or gelling agents and/or solvents. Such bases may thus, for example, include water and/or an oil such as liquid paraffin or a vegetable oil such as arachis oil or castor oil or a solvent such as a polyethylene glycol. Thickening agents which may be used include soft paraffin, aluminium stearate, cetostearyl alcohol, polyethylene glycols, microcrystalline wax and beeswax.

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Lotions may be formulated with an aqueous or oily base and will in general also contain one or more emulsifying agents, stabilising agents, dispersing agents, suspending agents or thickening agents.

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Powders for external application may be formed with the aid of any suitable powder base, for example, talc, lactose or starch. Drops may be formulated with an aqueous or non-aqueous base also comprising one or more dispersing agents, solubilising agents or suspending agents. Powder compositions for inhalation will preferably contain lactose. Spray compositions may be formulated, for example, as aqueous solutions or suspensions or as aerosols delivered from pressurised packs, with the use of a suitable propellant, e.g. dichlorodifluoromethane, trichlorofluoromethane, dichlorotetra-fluoroethane, 1,1,1,2,3,3,3-heptafluoropropane, 1,1,1,2-tetrafluoroethane, carbon dioxide or other suitable gas.

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Intranasal sprays may be formulated with aqueous or non-aqueous vehicles with the addition of agents such as thickening agents, buffer salts or acid or alkali to adjust the pH, isotonicity adjusting agents or anti-oxidants.

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Capsules and cartridges of for example gelatin, or blisters of for example laminated aluminium foil, for use in an inhaler or insufflator may be formulated containing a powder mix of a compound of the invention and a suitable powder base such as lactose or starch.

Solutions for inhalation by nebulation may be formulated with an aqueous vehicle with the addition of agents such as acid or alkali, buffer salts, isotonicity adjusting agents or antimicrobials. They may be sterilised by filtration or heating in an autoclave, or presented as a non-sterile product.

The pharmaceutical compositions according to the invention may also be used in combination with other therapeutic agents, for example anti-inflammatory agents (such as corticosteroids (eg. fluticasone propionate, beclomethasone dipropionate, mometasone

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furoate, triamcinolone acetonide or budesonide) or NSAIDs (eg. sodium cromoglyçate, nedocromil sodium, PDE-4 inhibitors, leukotriene antagonists, iNOS inhibitors, tryptase and elastase inhibitors, beta-2 integrin antagonists and adenosine 2a agonists)) or beta adrenergic agents (such as salmeterol, saibutamol, formoterol, fenoterol or terbutaline and salts thereof) or antilnfective agents (eg. antibiotics, antivirals).

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The invention thus provides, in a further aspect, a combination comprising a compound of formula (I) or a physiologically acceptable salt or solvate thereof together with another therapeutically active agent, for example an anti-inflammatory agent such as a corticosteroid, NSAID, beta adrenergic agent or an anti-infective agent. A pharmaceutical composition comprising a compound of formula (I) or a physiologically acceptable salt or solvate thereof In combination together with a long acting β₂ adrenergic receptor agonist (eg. salmeterol or a salt or solvate thereof such as salmeterol xinafoate) is of particular interest.

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The combination referred to above may conveniently be presented for use in the form of a pharmaceutical formulation and thus pharmaceutical formulations comprising a combination as defined above together with a physiologically acceptable diluent or carrier thereof represent a further aspect of the invention.

The individual components of such combinations may be administered either sequentially or simultaneously in separate or combined pharmaceutical formulations. Appropriate doses of known therapeutic agents will be readily appreciated by those skilled in the art.

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Compounds of the invention may conveniently be administered in amounts of, for example, 0.001 to 500mg/kg body weight, preferably 0.01 to 500mg/kg body weight, more preferably 0.01 to 100mg/kg body weight, 1 to 4 times daily. The precise dose will of course depend on the age and condition of the patient and the particular route of administration chosen.

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The compounds of the Invention have the advantage that they may be more efficacious, show greater selectivity (eg. In that they selectively antagonise $\alpha4$ integrins relative to $\beta2$ integrins such as LFA-1 or VLA-5 ($\alpha\nu\beta1$), have fewer side effects, have a longer duration of action, be less bloavailable or show less systemic activity when administered by inhalation, have ready and economic synthesis, or have other more desirable properties than similar known compounds.

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Certain intermediates are new and provide a further aspect of the invention.

The invention may be illustrated by reference to the following examples:

Examples

General Experimental Details

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is accelerated by an applied pressure of nitrogen at up to 5 p.s.i. Where thin fayer Where compounds were purified by "flash column chromatography on silica gel" this refer to the use of silica gel, 0.040 to 0.063mm mesh (e.g. Merck Art 9385), where column elutio chromatography (TLC) has been used this refers to silica gel TLC using 5 x 10 cm silica ge plates (e.g. Polygram SIL G/UV₂₅₄).

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Mass Spectrometry (MS) was carried out using an HP5989A Engine Mass Spectromete connected to a flow Inject system (0.05M aqueous ammonium acetate/methanol (35:65) at

flow rate of 0.7 ml/min) with positive thermospray ionisation. 15

NMR spectra were run on a Bruker DPX400 400MHz spectrometer.

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The Liquid Chromatography Mass Spectrometry (LCMS) system used was as follows: -

A 3µm ABZ+PLUS, 3.3cm x 4.6mm internal diameter column eluting with solvents: A –0.01M 0.05%v/v formic acid with a flow rate of 3ml/min. The following gradient protocol was used: 100% A for 0.7 mins; A+B mixtures, gradient profile 0 - 100% B over 3.7 mins; hold at 100% Aqueous ammonlum acetate + 0.1%v/v formic acid, and B - 95:5 acetonitrile/water

B for 0.9 mins; return to 0% B over 0.2 mins. 25

Positive and negative electrospray ionisation was employed

Protection Measurement

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- the resin was filtered. To 50µL of the filtrate was added 20% piperidine in DMF (0.95ml) an the absorbance of the solution was measured at 302nm using a UV spectrophotometer To 10mg of resin was added 20% piperidine in DMF (1ml). After shaking for 30 mins at 20° The method for measuring the substitution of Fmoc-amino acid resins was as follows:-Substitution was calculated using the following equation:-
- Substitution (mmol/g) = (Absorbance $\times 2 \times 10^4$) / (Extinction coefficient \times weight in mg) 35

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Intermediates

Intermediate 1: Methyl (2S)-2-(((2S)-2-((tert-butoxycarbonyl)amino]-4-methyl

pentanoyi}amino)-3-(4-hydroxyphenyl)propanoate

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a solution of N-(text-butoxycarbonyl)-L-leucine (7g) in acetonitrile (100ml), under a lyrosine methyl ester (5.5g) was added and stirring was continued for 18h. The mixture was concentrated in vacuo to ca. 10ml and the residue was partitioned between 1M hydrochloric acld (200ml) and ethyl acetate (100ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (100ml). The combined organic extracts were 1-(3-dimethylaminopropyl)-3-ethylcarbodilmide nydrochloride (5.9g) and 1-hydroxybenzotriazole (4.2g). After stirring for 30 mins at 20°C Lwashed with saturated aqueous sodium hydrogen carbonate (100ml), water (2 x 100ml) and evaporated with chloroform to give the title compound as a white foam (11.3g, 98%). LCMS: brine (50ml), dried over sodium sulphate and evaporated in vacuo. The residue was coadded was atmosphere,

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ntermediate 2: Methyl (2S)-2-{[(2S)-2-amino-4-methylpentanoyl]amino}-3-(4hydroxyphenyl)propanoate hydrochloride

R, 3.11 min; m/z 409 (MH*).

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To a solution of Intermediate 1 (3.1g) in 1,4-dioxane (10ml) was added 4M hydrogen vacuo. The residue was co-evaporated with toluene (2 x 20ml) and ether (2 x 20ml) to give chloride in 1,4-dioxane (20ml). The solution was stirred for 2h at 20°C then evaporated in the title compound as a white solid (2.6g, 98%). LCMS: R, 1.88 min; m/z 309 (MH+)

Intermediate 3: Methyl (2S)-3-(4-hydroxyphenyl)-2-(((2S)-4-methyl-2-[(2-{[3-(1-

- (0.31g) and 1-hydroxybenzotriazole (0.22g). After stirring for 30 mins at 20°C Intermediate 2 aqueous phase was further extracted with ethyl acetate (30ml). The combined organic (20ml), under a nitrogen added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.5g) was added followed by diisopropylethylamine (0.28ml) and stirring was continued for 18h. The mixture was concentrated in vacuo and the residue was partitioned between 2M hydrochloric acid (50ml) and ethyl acetate (30ml). The layers were separated and the piperidinylcarbonyl)-2-naphthyljoxy}acetyl)aminojpentanoyl}amlno)propanoate To a suspension of Intermediate 44 (0.45g) in acetonitrile afmosphere, was 25 30
- extracts were washed with saturated aqueous sodium hydrogen carbonate (30ml), water (2 x product was purified by flash column chromatography on silica gel eluting with ethyl 30ml) and brine (20ml), dried over sodium sulphate and evaporated in vacuo. The crude

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acetate/petroleum ether (2:1) to give the title compound as a white foam (0.6g, 69%). LCMS: R, 3.42 min; m/z 604 (MH*).

Intermediate 4: Methyl (2S)-3-(4-hydroxyphenyl)-2-[((2S)-2-{[2-(2-lodophenoxy)

acetyljamino}-4-methylpentanoyí)aminojpropanoate

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This was similarly prepared from Intermediate 43 (0.81g) and Intermediate 2 (1.02g). The crude product was purified by flash column chromatography on silica gel eluting with ethy acetate/cyclohexane (1:1) to give the title compound as a white foam (1.2g, 74%). LCMS: R. 3.40 min; m/z 569 (MH*).

Intermediate 5: Methyl (2S)-2-(((2S)-2-[(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-

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methylpentanoyl)amino)-3-(4-hydroxyphenyl)propanoate

This was similarly prepared from Intermediate 45 (0.29g) and Intermediate 2 (0.5g). The crude product was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (1:1) to give the title compound as a white foam (0.66g, 97%). LCMS: R, 3.55 min; m/z 503 (MH*).

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Intermediate 6: Methyl (2S)-2-({(2S)-2-[(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl)amino)-3-(4-{[(4-nitrophenoxy)carbonyl]oxy)phenyl)propanoate

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To a solution of intermediate 5 (0.59g) in dichloromethane (5ml), under a nitrogen atmosphere, was added 4-dimethylaminopyridine (0.18g). The mixture was cooled to 0-5°C and 4-nitrophenyl chloroformate (0.3g) was added. Stirring was continued for 18h allowing the reaction to warm to 20°C. The solution was diluted with chloroform (60ml) and washed with 1M hydrochloric acid (2 x 40ml) and water (40ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with cyclohexane/ethyl acetate (3:2) to give the title compound as a white foam (0.38g, 46%). LCMS: R₄ 3.98 min; *m/z* 668 (MH*).

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Intermediate 7: 4-[(2S)-2-(((2S)-2-((Tert-butoxycarbonyl)amino]-4-methyl pentanoyl)amino)-

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3-methoxy-3-oxopropyl]phenyl 4-[(2-phenylacetyl)amino]-1-piperidinecarboxylate

To a solution of triphosgene (0.59g) in anhydrous dichloromethane (40ml), under a nitrogen
atmosphere, was added a solution of Intermediate 1 (1.87g) in anhydrous dichloromethane
(10ml) followed by diisopropylethylamine (1.2ml). After stirring for 3h at 20°C Intermediate 59
(1g) was added followed by diisopropylethylamine (0.8ml). Stirring was continued for 18h
then the mixture was evaporated *in vacuo*. The crude product was purified by flash column

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chromatography on silica gel eluting with ethyl acetate/cyclohexane (1:1 swltching to 5:1) to give the title compound as a white solid (1.76g, 59%).

LCMS: R, 3.42 min; m/z 651 [M-H]

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Intermediate 8: 4-((2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-methoxy-3-oxopropyl)phenyl 4-[(2-phenylacetyl)amino]-1-piperidinecarboxylate hydrochloride To a solution of intermediate 7 (1.76g) in 1,4-dioxane (10ml) was added 4M hydrogen chloride in 1,4-dioxane (8ml). After stirring for 3h at 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a cream solid (1.59g, 100%). LCMS: R, 2.50 min; m/z 553 (MH*).

Intermediate 9: Methyl (2S)-2-({(2S)-2-{(tert-butoxycarbonyl)amino}-4-methyl pentanoyl)amino}-3-(4-{[(4-nitrophenoxy)carbonyl]oxy)phenyl)propanoate

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To a solution of Intermediate 1 (0.41g) in dichloromethane (3ml), under a nitrogen atmosphere, was added pyridine (1ml). The mixture was cooled to 0-5°C and 4-nitrophenyl chloroformate (0.22g) was added. Stirring was continued for 18h allowing the reaction to warm to 20°C. The solution was diluted with dichloromethane (40ml) and washed with 1M hydrochloric acid (50ml). The aqueous phase was further extracted with dichloromethane (40ml) and the combined organic extracts were dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with petroleum ether/ethyl acetate (3:1 switching to 3:2) to give thetitle compound as a white solld (0.29g, 50%). LCMS: R, 3.39 min; m/z 574 (MH7).

Intermediate 10: 4-((2S)-2-{[(2S)-2-Amino-4-methylpentanoyi]amino}-3-methoxy-3-

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oxopropyl)phenyl 4-[(2,2-dicyclohexylacetyl)amlno]-1-piperidinecarboxylate hydrochloride
To a solution of Intermediate 9 (0.22g) in anhydrous dichloromethane (4ml), under a nitrogen
atmosphere, was added Intermediate 58 (0.14g) followed by diisopropylethylamine (0.08ml).

After stirring for 4h at 20°C the mixture was diluted with dichloromethane (50ml), washed
with saturated aqueous potassium carbonate (3 x 25ml) and 1M hydrochloric acid (40ml),
dried over sodium sulphate and evaporated in vacuo to give a cream solid. To this was
added 4M hydrogen chloride in 1,4-dioxane (3ml) and the mixture was stirred for 3h at 20°C.
The solvent was evaporated in vacuo and the residue was triturated with ether to give the
title compound as a cream solld (0.24g, 95%). LCMS: R, 3.05 mln; m/z 641 (MH*).

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Intermediate 11: Tert-butyl (2S)-2-{((2S)-2-{((benzyloxy)carbonyl]amino}-4-methylpentanoyl)amino}-3-(4-hydroxyphenyl)propanoate

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To a solution of N-carbobenzyloxy-L-leucine (8.6g) in acetonitrile (150ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (6.83g) and 1-hydroxybenzotriazole (4.81g). After stirring for 30 mins at 20°C L-tyrosine*tert*-butyl ester (7.7g) was added and stirring was continued for 18h. The mixture was concentrated *in vacuo* to ca. 10ml and the residue was partitioned between 1M hydrochloric acid (300ml) and ethyl acetate (150ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (150ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (150ml), water (2 x 150ml) and brine (100ml), dried over sodium sulphate and evaporated *in vacuo*. The residue was coevaporated with chloroform to give the title compound as a white foam (15g, 96%). LCMS: R₁ 3.56 mln; *m/z* 485 (MH⁺).

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Intermediate 12: Tert-butyl (2S)-2-[((2S)-2-((benzyloxy)carbonyl)amino}-4-methyl pentanoyl)amino}-3-(4-(((4-nitrophenoxy)carbonyl]oxy}phenyl)propanoate

To a solution of Intermediate 11 (1.36g) in dichloromethane (15ml), under a nitrogen atmosphere, was added 4-nitrophenyl chloroformate (0.75g) and 4-dimethylaminopyridine (0.47g). The mixture was stirred for 18h at 20°C then diluted with chloroform (50ml), washed with 1M hydrochloric acid (2 x 30ml) and water (30ml), dried over sodium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with petroleum ether/ethyl acetate (4:1 switching to 1:1) to give the title compound as a white solid (1.34g, 74%). LCMS: R, 3.89 min; *m/z* 650 (MH*).

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Intermediate 13: 4-[(2S)-2-[((2S)-2-[((Benzyloxy)carbonyl]amino]-4-methyl pentanoyl)amino]-3-(tert-butoxy)-3-oxopropyl]phenyl 4-morpholinecarboxylate

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To a solution of intermediate 12 (0.34g) in dichloromethane (8ml), under a nitrogen atmosphere, was added morpholine (0.08ml) and disopropylethylamine (0.15ml). The mixture was stirred for 18h at 20°C then diluted with chloroform (30ml), washed with saturated aqueous potassium carbonate (3 x 40ml), 2M hydrochloric acid (40ml) and water (30ml), dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with ethyl acetate/petroleum ether (3:2) to give the title compound as a colourless gum (0.31g, 99%). LCMS: R, 3.60 min; m/z 598

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Intermediate 13 (Alternative procedure): 4-{(2S)-2-{((2S)-2-{((Benzyloxy) carbonyl]amino}-4-methylpentanoyl)amino}-3-{(tert-butoxy)-3-oxopropyl]phenyl 4-morpholinecarboxylate

To a solution of triphosgene (2.24g) in anhydrous dichloromethane (50ml), under a nitrogen atmosphere, was added a solution of Intermediate 11 (10g) in anhydrous THF (50ml) followed by dilsopropylethylamine (3.94ml). After stirring for 4h at 20°C morpholine (2ml) was added followed by dilsopropylethylamine (3.94ml). Stirring was continued for 18h then the mixture was partitioned between 1M hydrochloric acid (100ml) and ethyl acetate (75ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (75ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (50ml), water (50ml) and brine (30ml), dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with cyclohexane/ethyl acetate (3:1 switching to 1:1) to give the title compound as a white solid (6.89, 58%).

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Intermediate 14: 4-[(2S)-2-[((2S)-2-[((Benzyloxy)carbonyl]amino}-4-methyl pentanoyl)amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-(aminocarbonyl)-1-piperidinecarboxylate

This was similarly prepared from intermediate 11 (9g) and isonipecotamide (5.2g). The crude product was purified by flash column chromatography on silica gel eluting with ethyl acetate to give the title compound as a white solid (3.52g, 30%).

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Intermediate 14: (Alternative Procedure) 4-[(2S)-2-[((2S)-2-[((Benzyloxy) carbonyl]amino]-4-methyl pentanoyl)amino]-3-(tert-butoxy)-3-oxopropyl]phenyl 4-(aminocarbonyl)-1-piperidinecarboxylate

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To a solution of Intermediate 12 (1g) in dichloromethane (20ml), under a nitrogen atmosphere, was added isonipecotamide (0.23g) and dilsopropylethylamine (0.43ml). The mixture was stirred for 18h at 20°C then diluted with chloroform (80ml), washed with saturated aqueous potassium carbonate (3 x 50ml), 2M hydrochloric acid (50ml) and water (50ml), dried over sodium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with petroleum ether/ethyl acetate (3:2) switching to ethyl acetate/methanol (4:1) to give the title compound as a white solid (0.46g, 47%). LCMS: R, 3.47 min; *m*/z 639 (MH*).

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Intermediate 15: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-morpholinecarboxylate

To 10% palladium on carbon, Degussa type E101 (0.09g), under a nitrogen atmosphere, was added a solution of Intermediate 13 (0.3g) in ethanol (20ml) followed by ammonlum formate (0.17g). After stirring for 4h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Aid and the pad was washed with ethanol (10ml). The combined filtrate and washings were evaporated *in vacuo* and the residue was partitioned between dichloromethane (50ml) and 1M sodium hydroxide (15ml). The layers were separated and the organic phase was further washed with 1M sodium hydroxide (15ml) and water (15ml), dried over sodium sulphate and evaporated *in vacuo* to give the title compound as a grey gum (0.1g, 41%). LCMS: R, 2.43 min; *m/z* 464 (MH*).

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Intermediate 16: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-(aminocarbonyl)-1-piperidinecarboxylate

This was similarly prepared from intermediate 14 (0.46g). The title compound was obtained as a pale yellow gum (0.36g, 99%). LCMS: R, 2.33 min; m/z 505 (MH*).

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Intermediate 17: 4-[(2S)-2-[((2S)-2-[(Benzyloxy)carbonyl]amino}-4-methyl pentanoyl)amino]-3-(tert-butoxy)-3-oxopropyl]phenyl 4-acetyl-1-piperazine carboxylate

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To a solution of triphosgene (0.24g) in anhydrous dichloromethane (5ml), under a nitrogen atmosphere, was added a solution of Intermediate 11 (1g) in anhydrous THF (10ml) followed by diisopropylethylamine (0.43ml). After stirring for 4h at 20°C 1-acetylpiperazine (0.32g) was added followed by diisopropylethylamine (0.43ml). Stirring was continued for 18h then the mixture was partitioned between 1M hydrochloric acid (100ml) and ethyl acetate (75ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (75ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (50ml), water (50ml) and brine (30ml), dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with ethyl acetate switching to ethyl acetate/ethanol (9:1) to give the title compound as a white foam (1.3g, 99%). LCMS: R, 3.44 min; m/z 639 (MH*).

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Intermediate 18: 4-[(2S)-2-[((2S)-2-{[(Benzyloxy)carbonyl]amino}-4-methyl pentanoyl)amino]-3-(tert-butoxy)-3-oxopropyl]phenyl 4-benzoyl-1-piperazine carboxylate

To a solution of triphosgene (0.24g) in anhydrous dichloromethane (5ml), under a nitrogen atmosphere, was added a solution of Intermediate 11 (1g) in anhydrous THF (10ml) followed

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by diisopropylethylamlne (0.43ml). After stirring for 4h at 20°C Intermediate 56 (0.78g) was added followed by diisopropylethylamine (1.15ml). Stirring was continued for 18h then the mixture was partitioned between 1M hydrochloric acid (100ml) and ethyl acetate (75ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (75ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (50ml), water (50ml) and brine (30ml), dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with ethyl acetate/petroleum ether (1:1 switching to 2:1) to give the title compound as a white foam (1.02g, 71%). LCMS: R, 3.71 min; m/z 701 (MH*).

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Intermediate 19: 4-[(2S)-2-[((2S)-2-([(Benzyloxy)carbonyl]amino]-4-methyl pentanoyl)amino]-3-(tert-butoxy)-3-oxopropyl]phenyl 4-(1-piperidinylcarbonyl)-1-piperidinecarboxylate

This was similarly prepared from Intermediate 11 (1.81g) and Intermediate 55 (0.91g). The crude product was purified by flash column chromatography on sillca get eluting with

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dichloromethane/methanol (20:1) to give the title compound as a white foam (1.24g, 47%). LCMS: R, 3.63 min; m/z 707 (MH*).

Intermediate 20: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-(1-plperidlnylcarbonyl)-1-plperidinecarboxylate

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To 10% palladium on carbon, Degussa type E101 (0.27g), under a nitrogen atmosphere, was added a solution of Intermediate 19 (1.24g) in ethanol (20ml) followed by ammonium formate (0.77g). After stirring for 4h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Aid and the pad was washed with ethanol (20ml). The combined filtrate and washings were evaporated *in vacuo* and the residue was partitioned between dichloromethane (50ml) and 1M sodlum hydroxide (15ml). The layers were separated and the organic phase was further washed with 1M sodium hydroxide (15ml) and water (15ml), drled over sodium sulphate and evaporated *in vacuo* to give the title compound as a white foam (0.55g, 54%). LCMS: R, 2.63 min; *mz* 573 (MH²).

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30 Intermediate 21: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amlno}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-acetyl-1-plperazinecarboxylate hydrochloride

To 10% palladium on carbon, Degussa type E101 (0.4g), under a nitrogen atmosphere, was added a solution of Intermediate 17 (1.28g) in ethanol (30ml) followed by ammonium formate (0.38g). After stirring for 6h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Aid and the pad was washed with ethanol (20ml). The combined filtrate and washings

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were evaporated *in vacuo* and the residue was partitioned between dichloromethane (70ml) and 1M sodium hydroxide (30ml). The layers were separated and the aqueous phase was further extracted with dichloromethane (2 x 50ml). The combined organic extracts were dried over sodium sulphate. The solution was treated with 4M hydrogen chloride in 1,4-dioxane (0.55ml) and evaporated *in vacuo* to give the title compound as a white solid (1.02g, 94%). LCMS: R, 2.46 min; *m*/z 505 (MH*).

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Intermediate 22: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-benzoyl-1-piperazinecarboxylate hydrochloride

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To 10% palladium on carbon, Degussa type E101 (0.3g), under a nitrogen atmosphere, was added a solution of Intermediate 18 (1g) in ethanol (30ml) followed by ammonium formate (0.27g). After stirring for 6h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Ald and the pad was washed with ethanol (20ml). The combined filtrate and washings were evaporated *in vacuo* and the residue was partitioned between dichloromethane (70ml) and 1M sodium hydroxide (30ml). The layers were separated and the aqueous phase was further extracted with dichloromethane (2 x 50ml). The combined organic extracts were dried over sodium sulphate. The solution was treated with 4M hydrogen chloride in 1,4-dioxane (0.4ml) and evaporated *in vacuo* to give the title compound as a white solid (0.8g, 100%). LCMS: R, 2.72 min; *m*/z 567 (MH*).

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Intermediate 23: 4-[(2S)-2-{[(2S)-2-Amino-4-methylpentanoyl]amino}-3-(tert-butoxy)-3-oxopropyl]phenyl 4-morpholinecarboxylate hydrochloride

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To 10% palladium on carbon, Degussa type E101 (2.1g), under a nitrogen atmosphere, was added a solution of intermediate 13 (6.8g) in ethanol (500ml) followed by ammonium formate (4.1g). After stirring for 17h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Aid and the pad was washed with ethanol (50ml). The combined filtrate and washings were evaporated *in vacuo* and the residue was partitioned between dichloromethane (150ml) and 1M sodium hydroxide (75ml). The layers were separated and the aqueous phase was further extracted with dichloromethane (2 × 100ml). The combined organic extracts were dried over sodium sulphate. The solution was treated with 1M hydrogen chloride in ether (13ml) and evaporated *in vacuo*. The residue was triturated with ether to give the title compound as a white solid (4.8g, 87%).

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LCMS: R, 2.50 mln; m/z 464 (MH*).

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foam (2.1g, 94%). LCMS: R, 3.83 min; m/z 541 (MH*).

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Intermediate 24: 4-{(2S)-2-{((2S)-2-Amino-4-methylpentanoyi]amino}-3-(tert-butoxy)-3-oxopropyi]phenyl 4-(aminocarbonyl)-1-piperidinecarboxylate hydrochloride

To 10% palladium on carbon, Degussa type E101 (1.1g), under a nitrogen atmosphere, was added a solution of intermediate 14 (3.41g) in ethanol (80ml) followed by ammonium formate (2.1g). After stirring for 3h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Ald and the pad was washed with ethanol (40ml). The combined filtrate and washings were evaporated in vacuo and the residue was partitioned between chloroform (500ml) and saturated aqueous sodium hydrogen carbonate (2 × 100ml). The layers were separated and the aqueous phase was further extracted with chloroform (2 × 100ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (3 × 100ml) and water (2 × 100ml) then dried over sodium sulphate. The solution was treated with 4M hydrogen chloride in 1,4-dioxane (1.5ml) and evaporated in vacuo. The residue was azeotroped with toluene (2 × 50ml) to give the title compound as a white solid (2.88g, 100%). LCMS: R, 2.36 min; m/z 505 (MH*).

Intermediate 25: Tert-butyl (2S)-2-([(2S)-2-((2-[2-(tert-butyl)phenoxy]acetyl) amino)-4-methylpentanoyl]amino)-3-(4-hydroxyphenyl)propanoate

To 10% palladium on carbon, Degussa type E101 (0.63g), under a nitrogen atmosphere, was added a solution of Intermediate 11 (2g) in ethanol (20ml) followed by ammonium formate (1.8g). After stirring for 2h at 20°C the mixture was filtered through a pad of Harborlite J2 Filter Ald and the pad was washed with ethanol (50ml). The combined filtrate and washings were evaporated in vacuo and the residue was partitioned between dichloromethane (100ml) and saturated aqueous sodium hydrogen carbonate (50ml). The layers were separated and the organic phase was further washed with saturated aqueous sodium hydrogen carbonate (50ml) and water (50ml), dried over magnesium sulphate and evaporated in vacuo to give a white solid. A solution of this in DMF (5ml) was added to a premixed solution of Intermediate 46 (0.87gg), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.80gg) and 1-hydroxybenzothazole (0.57gg) in acetonitrile (10ml) which had been stirring under a nitrogen atmosphere for 30 mins at 20°C. Stirring was continued for 18h. The mixture was diluted with ethyl acetate (200ml), washed with 1M hydrochloric acid (3 × 50ml), saturated aqueous sodium hydrogen carbonate (3 × 50ml) and brine (50ml), dried over magneslum sulphate and evaporated in vacuo to give the title compound as a white

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Intermediate 26: Tert-butyl (2S)-2-{((2-(2-(tert-butyl)phenoxy)acetyl) amino)-4-

methylpentanoyljamino}-3-(4-{[(4-nitrophenoxy)carbonyijoxy)phenyl) propanoate

To a solution of Intermediate 25 (2.1g) in dichloromethane (20ml); under a nitrogen atmosphere, was added 4-nitrophenyl chloroformate (1.1g) and 4-dimethylaminopyridine (0.69g). The mixture was stirred for 18h at 20°C then diluted with chlorofdrm (80ml), washed with 1M hydrochloric acid (2 x 50ml) and water (50ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with cyclohexane/ethyl acetate (2:1) to give thetitle compound as a clear oil (2.65g, 97%). LCMS: R, 4.17 min; *m/*z 708 (MH*).

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Intermediate 27: 4-[(2S)-2-((2S)-2-[(2-Bromoacetyl)amino]-4-methylpentanoyl} amino)-3-(tert-butoxy)-3-oxopropyl]phenyl 4-morpholinecarboxylate

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A solution of Intermediate 23 (0.5g) and diisopropylethylamine (0.19ml) in dichloromethane (10ml) was cooled to 0.5°C. To this was added bromoacetyl chloride (0.09ml) followed by diisopropylethylamine (0.19ml) and stirring was continued for 2h. The mixture was diluted with dichloromethane (50ml), washed with 2M hydrochloric acid (50ml), saturated aqueous sodium hydrogen carbonate (50ml) and brine (30ml), dried over magnesium sulphate and evaporated in vacuo to give the title compound as a white foam (0.52g, 89%).

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LCMS: R, 3.28 min; m/z 584 (MH*).

Intermediate 28:·4-[(2S)-2-(((2S)-2-[(2-Bromoacetyl)amino]-4-methylpentanoyl) amino)-3-methoxy-3-oxopropyl]phenyl 4-[(2-phenylacetyl)amino]-1-piperidine carboxylate

To a solution of Intermediate 8 (0.48g) in anhydrous dichloromethane (4ml) was added disopropylethylamine (0.142ml). The mixture was cooled to 0-5°C and bromoacetyl chloride (0.07ml) was added. Stirring was continued for 1h allowing the reaction to warm to 20°C. The mixture was diluted with dichloromethane (5ml) and washed with saturated aqueous sodium hydrogen carbonate (5ml), water (10ml) and brine (10ml), dried over sodium sulphate and evaporated *in vacuo* to give the title compound as a white solid (0.464g, 85%). LCMS: R, 3.20 min; *mix* 672 [M-H].

Intermediate 29: (2S)-3-[4-(Allyloxy)phenyl]-2-([(9H-fluoren-9-ylmethoxy)

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carbonyl]amino)propanoic acid bound to Wang resin via acid

To Wang resin (100-200 mesh, 10g) was added a solution of (2S)-3-[4-(aliyloxy)phenyl]-2-{[(9H-fluoren-9-ylmethoxy)carbonyl]amino}propanoic acid (8.5g) in DMF (45ml). After 15 mins pyridine (2.4ml) was added followed by 2,6-dichlorobenzoyl chloride (2.75ml). The

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mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 40ml), dichloromethane (5 x 40ml) and ether (5 x 40ml) then dried in vacuo. The amount of (2S)-3-[4-(aliyloxy)phenyl]-2-[(9H-fluoren-9-ylmethoxy)carbonyl]amino} propanolc acid substituted on the resin was calculated to be 0.52 mmol/g.

Intermediate 30: (2S)-3-[4-(Allyloxy)phenyl]-2-[((2S)-2-[[(9H-fluoren-9-ylmethoxy)

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carbonyljamino}-4-methylpentanoyl)amino]propanoic acid bound to Wang resin via acid Intermediate 29 (2.5mmol) was treated with 20% piperidine in DMF (15ml) and shaken for 1h 30mins at 20°C. The resin was filtered and washed with DMF (5 x 20ml). A solution of Fmocleucine (2.8g) in DMF (10ml) was added followed by a solution of benzotriazol-1-yl-oxytrispyrrolidinophosphonium hexafluoro phosphate (4.1g) in DMF (5ml) and diisopropylethylamine (2.8ml). The mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 20ml), dichloromethane (5 x 20ml) and ether (5 x 20ml) then dried *in vacuo*. A 5mg sample was treated with trifluoroacetic acid/ dichloromethane

Intermediate 31: (2S)-3-[4-(Allyloxy)phenyl]-2-[[(2S)-2-((2-[2-(tert-butyl)phenoxy] acetyl)amino)-4-methylpentanoyl]amino)propanoic acid bound to Wang resin via acid

(1:1) (1ml) for 0.5h at 20°C, resin was filtered and the filtrate analysed by LCMS: R, 4.22 min;

m/z 557 (MH*).

- at 20°C. The resin was filtered and washed with DMF (5 x 10ml). A solution of Intermediate 48 (0.314g) in DMF (10ml) was added followed by a solution of benzotriazol-1-yl-oxytrispyrrolidinophosphonium hexafluoro phosphate (0.78g) in DMF (5ml) and disopropylethylamine (0.68ml). The mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 10ml), dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried *in vacuo*. A 5mg sample was treated with trifluoroacetic acid/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was filtered and the filtrate analysed by LCMS: R, 4.27 min; mtz 525 (MH*).
- Intermediate 32: (2S)-3-{4-(Allyloxy)phenyl]-2-[((2S)-4-methyl-2-{[2-(2-methyl-2-(2-methyl

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Intermediate 33: (2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy)acetyl}amino)-4-

methylpentanoyi]amino}-3-(4-{[(4-nitrophenoxy)carbonyi]oxy}phenyl)propanoic acid bound to Wang resin via acid

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(0.1g). The mixture was shaken for 40 mins at 20°C. The resin was filtered and washed with dichtoromethane (5 x 10ml) then retreated with a solution of phenylsilane (1ml) i dichloromethane (9ml) followed by tetrakis(triphenylphosphlne)palladium(0) (0.1g). Afte of dilsopropylethylamine (1.74ml) in 1: dichloromethane/THF (16ml). 4-Nitrophenyl chloroformate (2g) was added portionwise and the mixture was shaken for 18h at 20°C. The resin was filtered and washed with dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried *in vacuo.* A 5mg sample was Ireated with trifluoroacetic acld/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was shaking for 40 mins at 20°C the resin was filtered and washed with dichloromethane (5 phenylsllane (1ml) dichloromethane (9ml) followed by tetrakls(triphenytphosphine)palladium(0) filtered and the filtrate analysed by LCMS: R, 4.33 min; m/z 650 (MH*). Intermediate 31 (1mmol) was treated with a solution of a solution treated with then

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Intermediate 34: (2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino}
pentanovi)aminol-3-(4-{[(4-nitrophenoxy)carbonylloxy)phenylloropanoic acid hound

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pentanoyl)amino]-3-(4-[[(4-nitrophenoxy)carbonyl]oxy)phenyl)propanoic acid bound to Wang resin via acid

This was similarly prepared from Intermediate 32 (0.97mmol). LCMS: R_t 3.31 min; *m/z* 443 (MH*).

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Intermediate 35: (2S)-2-[((2S)-2-[[(9H-Fluoren-9-ylmethoxy)carbonyl]amino}-4-

methylpentanoyl)amino]-3-(4-[[(4-nitrophenoxy)carbonyl]oxy)phenyl)propanoic acid bound to Wang resin via acid

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This was similarly prepared from intermediate 30 (1.05mmol). LCMS: R₁ 4.32 min; *m/z* 682 (MH*).

Intermediate 36: (2S)-2-[((2S)-2-{((9H-Fluoren-9-y/methoxy)carbonyl]amino}-4-

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methylpentanoyl)aminoj-3-[4-({[4-(2-furoyl)-1-piperazinyl]carbonyl}oxy)phenyl] propanoic acid bound to Wang resin via acid

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Intermediate 35 (1.05mmol) was treated with a solution of 1-(2-furoyl)piperazine (0.57g) in 1:1 dichloromethane/THF (9ml) followed by diisopropylethylamine (1.1ml). After shaking for 4h at 20°C the resin was filtered and washed with dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried in vacuo. A 5mg sample was treated with trifluoroacetic acid/

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dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was filtered and the filtrate analysed by LCMS: R, 3.67 min; m/z 723 (MH*).

Intermediate 37: (2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amlno}-1-piperidinyl) carbonyl]oxy}phenyl)-2-[((2S)-2-{[(9H-fluoren-8-ylmethoxy)carbonyl]amino}-4-methylpentanoyl)amino]propanoic acid bound to Wang resin via acid

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This was similarly prepared from intermediate 35 (1.7mmol) and intermediate 53 (1.02g). LCMS: R, 4.03 min; m/z 795 (MH*).

Intermediate 38: (2S)-2-({(2S)-2-[(2-Bromoacetyl)amlno]-4-methylpentanoyl} amino)-3-[4-([4-(2-furoyl)-1-piperazinyl]carbonyl)oxy)phenyl]propanoic acid bound to Wang resin via acid

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Intermediate 36 (1.05mmol) was treated with 20% plperidine in DMF (8ml) and shaken for 1h 30mlns at 20°C. The resin was filtered and washed with DMF (5 x 10ml). A solution of bromoacetic acid (0.44g) in DMF (8ml) was added followed by 1,3-dlisopropylcarbodiimide (0.49ml). The mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 10ml), dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried in vacuo. A 5mg sample was treated, with trifluoroacetic acid/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was filtered and the filtrate analysed by LCMS: R, 3.11 min; m/z 621 (MH*).

Intermediate 39: (2S)-2-(((2S)-2-f((2-Bromoacetyl)amino]-4-methylpentanoyl) amino)-3-(4[((4-f((2-f((4-chlorophenyl)acetyl)amino)-1-piperidinyl)carbonyl)oxy) phenyl)propanoic acid
bound to Wang resin via acid

This was similarly prepared from Intermediate 37. (0.73mmol). LCMS: R₁ 3.43 min; m/z 695 (MH⁺).

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Intermediate 40: (2S)-3-[4-(Allyloxy)phenyl]-2-(((2S)-2-[(2-bromoacetyl)amino]-4-methylpentanoyl)amino)propanoic acid bound to Wang resin via acid

at 20°C. The resin was filtered and washed with 20% piperidine in DMF (6ml) and shaken for 1h at 20°C. The resin was filtered and washed with DMF (5 x 10ml). A solution of bromoacetic acid (0.23g) in DMF (3ml) was added followed by 1,3-dilsopropylcarbodiimide (0.26ml). The mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 10ml), dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried *in vacuo*. A 5mg sample was treated with trifluoroacetic acid/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was filtered and the filtrate analysed by LCMS: R, 3:47 min; *m*/z 455 (MH*).

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Intermediate 40 (0.55mmol) was treated with DMF (4ml). 2-Cyclohexylphenol (0.97g) potassium carbonate (0.76g) and sodium todide (0.82g) were added and the mixture was shaken for 40h at 20°C. The resin was filtered and washed with water (3 imes 5ml), DMF (5 imes5ml), dichloromethane (5 x 5ml) and ether (5 x 5ml) then dried in vacuo. A 5mg sample was treated with trifluoroacetic acid/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was acetyljamino)-4-methylpentanoyl)aminojpropanoic acid bound to Wang resin via acid Intermediate 41: (2S)-3-[4-(Allyloxy)phenyl]-2-[((2S)-2-{[2-(2-cyclohexylphenoxy) filtered and the filtrate analysed by LCMS: R, 4.49 min; m/z 551 (MH*).

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Intermediate 42: (2S)-2-[((2S)-2-[[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-

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methylpentanoyl)amlno]-3-(4-{[(4-nitrophenoxy)carbonyl]oxy}phenyl)propanoic acid bound to

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Intermediate 41 (0.55mmol) was treated with a solution of phenylsilane (1.35ml) In dichtoromethane (10ml) followed by tetrakis(triphenylphosphine)palladium(0) (0.063g). The mixture was shaken for 40 mins at 20°C. The resin was filtered and washed with dichloromethane (5 x 10ml) then retreated with a solution of phenylsilane (1.35ml) in dichloromethane (10ml) followed by tetrakis (triphenylphosphine) palladium (0) (0.063g). After shaking for 40 mins at 20°C the resin was filtered and washed with dichloromethane (5 x - dichloromethane/THF (8ml). 4-Nitrophenyl chloroformate (2.2g) was added portionwise and the mixture was shaken for 18h at 20°C. The resin was filtered and washed with dichloromethane (5 x 10ml) and ether (5 x 10ml) then dried in vacuo. A 5mg sample was treated with trifluoroacetic acid/ dichloromethane (1:1) (1ml) for 0.5h at 20°C, resin was .⊑ (1.9ml) diisopropylethylamine filtered and the filtrate analysed by LCMS: Rt 4.54 min; m/z 676 (MH*). ŏ solution ര with treated

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Intermediate 43: (2-lodophenoxy)acetic acid

fert-Butyl bromoacetate (4.0ml) was added to a suspension containing 2-lodophenol (4.98g) and potassium carbonate (6.3g) in DMF (40ml). The mixture was stirred for 1h at 20°C under a nitrogen atmosphere and was then partitioned between ethyl acetate (150ml) and water (100ml). The aqueous layer was extracted with fresh ethyl acetate (2 \times 80ml) and the combined organic extracts washed with brine (100ml), dried over magnesium sulphate and evaporated in vacuo to give a clear liquid (7.58g). This was dissolved in dichloromethane (20ml) and trifluoroacetic acld (8ml) and the solution stirred for 2h at 20°C. Solvent was evaporated in vacuo and the residue triturated in a mixture of cyclohexane/ethyl acetate

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277 (5:1) to give the title compound as a white solid (5.19g, 82%). LCMS: R, 3.02 min; m/z

Intermediate 44: {[3-(1-Piperidinylcarbonyl)-2-naphthylloxy]acetic acid

1977)((4.98g). The intermediate ester was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (1·1) and the title compound was isolated as This was similarly prepared from 3-(1-piperidinylcarbonyl)-2-naphthot (Griffiths and Hawkins, a white solid (3.2g, 53%). LCMS: R, 3.74 mln; m/z 314 (MH*).

Intermediate 45: Dibenzo[b,d]furan-4-carboxyllc acid 9

A solution of 1.6M n-butyllithium in hexane (18.5ml) was added dropwise to a stirred solution resulting suspension was allowed to warm to 20°C where it was stirred for 3h. It was then (250ml) under a nitrogen atmosphere. The resulting white suspension was allowed to stand for 1h at 20°C and was then diluted with 2M sodium hydroxide (500ml). The aqueous extract was washed with ether (3 x 200ml), acidified to pH 1 with 6M hydrochloric acid and extracted with ethyl acetate (3 x 200ml). The combined organic extracts were washed with brine cooled to -78°C and added to a mixture of excess solid carbon dioxide in diethyl ether (50ml), dried over magnesium sulphate and evaporated in vacuo to give the title compound of dibenzofuran (5.0g) in anhydrous THF (25ml) at -78°C under a nitrogen atmosphere. as a white solid (3.64g, 58%). LCMS: R, 5.06 mln; m/z 213 (MH*).

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Intermediate 46: [2-(Tert-butyl)phenoxy]acetic acid

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Methyl bromoacetate (3.0ml) was added to a suspension containing 2-fert-butylphenol (5.0ml) and potassium carbonate (10.6g) in DMF (250ml). The mixture was stirred for 20h at vas extracted with more ether (100ml) and the combined organic extracts washed with brine (1:9) to give a clear liquid (6.64g). This was dissolved in methanol (100ml) and 2M sodium partitioned between ether (200ml) and 1M hydrochloric acid (100ml). The aqueous layer (100ml), dried over magnesium sulphate and evaporated in vacuo. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane 20°C under a nitrogen atmosphere and was then evaporated in vacuo to a slurry which was evaporated in vacuo and the aqueous residue was washed with diethyl ether (50ml), combined organic extracts were washed with brine (50ml), dried over magnesium sulphate acidified to pH 1 with θ M hydrochloric acid and extracted with ethyl acetate (2 x 200ml). hydroxide (100ml) and the solution was stirred for 0.5h at 20°C.

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and evaporated *in vacuo* to give the <u>title compound</u> as a white crystalline mass (5.86g, 95% LCMS: R, 3.78 min; m/z 207 [M-H];

Intermediate 47: 4-(2-Methoxy-2-oxoethoxy)benzoic acid

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hydroxybenzoate (Shah et al., 1992) (3.03g), sodium iodide (2.55g) and potassiun nitrogen atmosphere and then allowed to cool to 20°C. It was then partitioned between water (50ml) and ethyl acetate (100ml) and the organic extract washed with water (2 imes 80ml and brine (60ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with a gradient o ethyl acetate/petroleum ether (1:9) to ethyl acetate/petroleum ether (1:2) to give a pale rec gum (3.85g). This was dissolved in dichloromethane (50ml) and trifluoroacetic acid (15ml) was added and the solution was stirred for 3h at 20°C. Solvents were evaporated in vacuo to give the title compound as a white solid (2.97g, 91%). LCMS: R, 2.45 min; m/z 211 (MH*). carbonate (4.2g) In acetonitrile (60ml). The mixture was stirred for 17h at 90°C under Methyl bromoacetate (1.6ml) was added to a suspension containing tert-butyl

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Intermediate 48: [4-(1-Piperidinylcarbonyl)phenoxy]acetic acid

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To a suspension of Intermediate 47 (2.95g) in acetonitrile (55ml) was added diisopropylethylamine (3.5ml) followed by (1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate (4.59). The resulting solution was stirred for 10mins at 20°C under a nitrogen atmosphere and then piperidine (1.4ml) was added and the mixture was stirred for 18h at 20°C under a nitrogen atmosphere and then evaporated in vacuo. The residue was partitioned between ethyl acetate (100ml) and 8% aqueous sodium hydrogen carbonate (65ml) and the organic extract was washed with 2M hydrochloric acid (50ml) and brine (100ml), dried over magnesium sulphate and evaporated in vacuo to give an orange oil (4.05g). This was dissolved in methanol (100ml) and 1M sodium hydroxide (30ml) was added and the mixture stirred for 3h at 20°C. It was then acidified to pH 1 with 1M hydrochloric acid and cooled to 5°C and the precipitate collected by filtration and dried in vacuo to give the title compound as a white solid (3.03g, 80%). LCMS: R, 4.17 min; m/z 264

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Intermediate 49: (2-Benzoylphenoxy) acetic acid

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hydroxybenzophenone (2.3g), potasslum carbonate (3.2g) and sodium iodide (2.33g) in acetonitrile (35ml). The mixture was stirred for 18h at 90°C under a nitrogen atmosphere and containing suspension æ ₽ added was (3.0ml)bromoacetate

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was then allowed to cool to 20°C. It was then partitioned between ethyl acetate (80ml) and water (60ml) and the organic extract washed with water (2 x 60ml) and brine (60ml), dried column chromatography on silica gel eluting with ethyl acetate/petroleum ether (1:1) to give over magnesium sulphate and evaporated in vacuo. The crude material was purified by flash a pale yellow oil (3.05g). This was dissolved in methanol (100ml) and 1M sodium hydroxide (35ml) and the solution was stirred for 18h at 20°C. The solution was acidified to pH 1 with 2M hydrochloric acid and extracted with ethyl acetate (2 x 80ml). The combined organic eluting with a gradient of ethyl acetate/petroleum ether (1:1) to ethyl acetate/methanol (4:1) extracts were washed with water (2 x 70ml), dried over magnesium sulphate and evaporated In vacuo. The crude product was purified by flash column chromatography on silica gel to give the title compound as a pale yellow gum (1.62g, 57%). LCMS: R, 3.41 min; m/z 257

ntermediate 50: [(1-Bromo-2-naphthyl)oxy]acetic acid

(1:3) and the title compound was isolated as a pale brown solid (11.36g, 89%). LCMS: R, This was similarly prepared from 1-bromo-2-naphthol (10.55g). The intermediate ester was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane 4.17 min; m/z 281 [M-H];

Intermediate 51: [4-(Aminocarbonyl)phenoxyjacetic acid 20

98% formic acid (50ml) was stirred under reflux for 2h and then cooled in an ice bath. The A solution of 4-formylphenoxyacetic acid (1.86g) and hydroxylamine hydrochloride (1.07g) in The mixture was diluted with water (100ml), washed with ethyl acetate (50ml) and acidified to pH 2 with 6M hydrochloric acid. The precipitate was collected by filtration, washed with precipitate was collected by filtration, washed with water and dried in vacuo to give a white (50ml) was stirred under reflux under a nitrogen atmosphere for 4h and then allowed to coof. water and dried in vacuo to give the title compound as a white solid (1.08g, 53%). LCMS: R A mixture of this with powdered potassium hydroxide (2.3g) in tert-butanol 1.90 min; m/z 196 (MH*). solld (1.1g).

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Intermediate 52: Tert-butyl 4-amino-1-piperidinecarboxylate

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Sodium triacetoxyborohydride (30.2g) was added portionwise over 10min to an ice-cooled mixture of 1-(lert-butoxycarbonyl)-4-piperidone (20.07g), dibenzylamine. (19.7g) and acetic solution was then treated cautiously with 2M sodium hydroxide (400ml) and the separated acid (5ml) in dichloromethane (500ml) and stirring was then continued for 16h at 20°C.

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organic layer, dried over magnesium sulphate and evaporated *In vacuo*. The residue was triturated in hexane/ether (2:1) (250ml) to give a white solid (18.75g). This was dissolved in a mixture of THF (50ml), ethanol (50ml) and 2M hydrochloric acid (8ml) and the solution added to a suspension of 20% palladium hydroxide on carbon (5.0g) in ethanol (100ml). The mixture was hydrogenated at 20°C and 1 etmosphere for 17h and was then filtered through a pad of Harborlite J2 Filter Aid and the pad washed with ethanol (100ml). The combined filtrate and washings were evaporated *in vacuo* and the residue dissolved in water (50ml) and adjusted to pH 9 with 2M sodium hydroxide and evaporated *in vacuo*. The residue was leached into a mixture of ethanol (30ml) and chloroform (70ml) and insoluble material removed by filtration. The mother liquors were evaporated *in vacuo* to give the title compound as a colourless oil (10.04g, 49%). LCMS: R, 1.81 mln; *m/z* 201 (MH+*).

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Intermediate 53: 2-(4-Chlorophenyl)-N-(4-piperidinyl)acetamide hydrochloride

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To a solution of 4-chlorophenylacetic acid (2.55g) in acetonitrile (100ml), under a nitroge added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (3.16g) and 1-hydroxybenzotriazole (2.22g). After stirring for 10 mins at 20°C a solution of Intermediate 52 (3g) in acetonitrile (20ml) was added, and stirring was continued for 18t The mixture was evaporated in vacuo and the residue partitioned between water (100ml and ethyl acetate (100ml). The organic phase was washed with saturated aqueous sodiur hydrogen carbonate (2 \times 80ml) and water (50ml), dried over magnesium sulphate an evaporated in vacuo to give a pale yellow solid. This was triturated with ether to give a whit A portion of this (2.36g) was dissolved in 1,4-dioxane (100ml) and 4h hydrogen chioride in 1,4-dioxane (12ml) was added. The solution was stirred for 18h s 20°C and then a further portion of 4M hydrogen chloride in 1,4-dioxan (8ml) was addec Stirring was continued for a further 18h at 20°C and the solution was evaporated in vacuo to give a white solld. This was triturated in ether to give the title compound as a white solk (1.9g, 77%). LCMS: R, 1.89 min; m/z 253 (MH*). was atmosphere,

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Intermediate 54: N-(4-Fluorobenzyl)-4-plperidinecarboxamide hydrochloride

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To a solution of 1-*tert*-butoxycarbonytplperidine-4-carboxylic acid (3.61g) in acetonitrile (25ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (3.21g) and 1-hydroxybenzotriazole (2.29g). After stirring for 20 mins at 20°C 4-fluorobenzylamine (2.0ml) was added and stirring was continued for 3h. The mixture was concentrated *in vacuo* and the residue was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (200ml). The layers were separated and the

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organic phase was washed with 1M hydrochloric acid (3 x 50ml), saturated aqueous sodium hydrogen carbonate (3 x 50ml) and brine (50ml), dried over magnesium sulphate and evaporated *In vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with a gradient of cyclohexane/ethyl acetate (1:1) to neat ethyl acetate to give colourless crystals (5.02g). A portion of this (4.96g) was dissolved in 1,4-dioxane (20ml) and 4M hydrogen chloride in 1,4-dioxane (15ml) was added. The mixture was stirred for 2h at 20°C and the precipitate was collected by filtration, washed with 1,4-dioxane and diethyl ether and dried *In vacuo* to give the title compound as a white hygroscopic solid (3.54g, 83%). LCMS: R, 1.52 min; *m/z* 237 (MH*).

Intermediate 55: 1-(4-Piperidinylcarbonyl)piperidine hydrochloride

This was similarly prepared from 1-tert-butoxycarbonylpiperidine-4-carboxylic acid (3.68g) and piperidine (1.6ml). The Intermediate amide was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol (10:1) and the title compound was isolated as a white solid (3.26g, 93%). MS: *m/z* 197 (MH*), TLC: R, 0.1 [dichloromethane/ethanol/880 ammonia (50:8:1) visualisation with iodoplatinic acid).

Intermediate 56: 1-Benzoylpiperazine

This was similarly prepared from benzolc acid (5.02g) and 1-(*tert*-butoxycarbonyl)piperazine (7.66g) and the title compound was isolated as a white solid (7.7g, 82%). LCMS: R, 0.51 min; m/z 191/MH*)

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Intermediate 57: 2-Cyclohexyl-N-(4-plperidinyl)acetamide

A solution of 4-amino-1-benzylpiperidine (5.0ml), cyclohexaneacetic acid (3.79g) and (1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroborate (8.35g) in acetonitrile (60ml) was stirred for 18h at 20°C under a nitrogen atmosphere and was then evaporated *in vacuo* to a syrup. This was partitioned between ethyl acetate (200ml) and saturated aqueous sodium hydrogen carbonate (2 x 100ml). The organic extract was washed with saturated aqueous sodium hydrogen carbonate (2 x 100ml) and brine (100ml), dried over magnesium sulphate and evaporated *in vacuo* to give an off-white solid. This was crystallised from cyclohexane to give cream crystals (6.24g). A portion of this (3.8g) was dissolved in ethanol (100ml) and treated with 10% paliadium on carbon, Degussa type E101 (1.2g) and ammonium formate (2.24g). The mixture was stirred for 2.5h at 20°C under a nitrogen atmosphere and was then filtered through a pad of Harborlite J2 Filter Aid and the pad washed with ethanol (100ml). The combined filtrate and washings were evaporated *in vacuo*

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and the residue was partitioned between chloroform (100ml) and 0.5M potassium hydroxide (10ml). The layers were separated and the aqueous phase extracted with fresh chloroform (2 x 100ml) and the combined organic extracts dried over magnesium suiphate and This was triturated with ether to give the title evaporated in vacuo to give a white solid. compound as a white solid (2.01g, 60%).

LCMS: R₁ 1.93 min; m/z 225 (MH*).

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Intermediate 58: 2,2-Dicyclohexyl-N-(4-piperidinyl)acetamide

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DMF A solution containing dicyclohexylacetic acid (4.75g), diisopropylethylamine (7.5ml) and (250ml) was stirred for 10min at 20°C and then 4-amino-1-benzylpiperidine (4.3ml) was added dropwise over 10min. The mixture was stirred for 18h at 20°C and was then diluted with ethyl acetate (200ml) and the precipitate collected by filtration, washed with ethyl acetate (60ml) and water (50ml) and dried *in vacuo* to give a white solid (5.91g). A portion of this (3g) was suspended in ethanol (300ml) and treated with 10% palladium on carbon, Degussa type E101 (1.2g) and ammonium formate (2.68g). The mixture was stirred for 4h at 20°C under a nitrogen atmosphere and was then filtered through a pad of Harborlite J2 Filter Aid and the pad washed with ethanol (50ml). The combined filtrate and washings were evaporated in vacuo and the residue was partitioned between chloroform (200ml) and 0.5M sodium hydroxide (150ml). The layers were separated and the aqueous phase extracted with fresh chloroform (100ml) and the combined organic extracts dried over magnesium sulphate and evaporated in vacuo to give a white solid. This was triturated with ice-cold ether to give the title compound as a white solid (1.8, 78%). LCMS: R, 2.69 min; m/z 307 phosphate (11g) benzotnazol-1-yl-oxy-trispyrrolidinophosphonium hexafluoro

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Intermediate 59: 2-Phenyl-N-(4-piperidinyl)acetamide

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a solution of phenylacetic acid (3.4g) in acetonitrile (100ml), under a nitrogen added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (5.28g) and 1-hydroxybenzotriazole (3.72g). After stirring for 30 mins at 20°C 4-amino-1benzylplperidine (5.1ml) was added and stirring was continued for 18h. The mixture was concentrated in vacuo and the residue was partitioned between 2M hydrochloric acid (100ml) and ethyl acetate (75ml). The layers were separated and the aqueous phase was washed with more ethyl acetate (75ml), basified with solid potassium carbonate and extracted with dichloromethane (2 x 100ml). The combined organic extracts were washed with water (2 x 100ml) and brine (50ml), dried over sodium sulphate and evaporated in atmosphere, was

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vacuo to give a white solid (4.8g). A portion of this (4.7g) was dissolved in ethanol (150ml) formate (2.88g). The mixture was stirred for 4h at 20°C under a nitrogen atmosphere and and treated with 10% palladium on carbon, Degussa type E101 (1.5g) and ammonium was then filtered through a pad of Harborlite J2 Filter Aid and the pad washed with ethanol partitioned between chloroform (100ml) and 0.5M sodium hydroxide (50ml). The layers were separated and the aqueous phase extracted with fresh chloroform (2 imes 100ml) and the combined organic extracts dried over sodium sulphate and evaporated in vacuo to give the (150ml). The combined filtrate and washings were evaporated in vacuo and the residue was 219 (MH*), dichloromethane/methanol/880 ammonia (40:10:1) visualisation with iodine], solid (2.4g, 45%). MS: m/z a white compound

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Example 1: (2S)-2-[((2S)-2-{[2-(2-Benzoylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[({4-[(2-phenylacetyl)amino]-1-piperidinyl}carbonyl) oxy]phenyl}propanolc acid

odide (0.1g). After stirring for 18h at 20°C the mixture was partitioned between saturated To a solution of 2-hydroxybenzophenone (0.134g) in anhydrous DMF (0.5ml) was added separated and the aqueous phase was further extracted with ethyl acetate (3 imes 10ml). The anhydrous potassium carbonate (0.093g) followed by Intermediate 28 (0.152g) and sodium aqueous sodium hydrogen carbonate (10ml) and ethyl acetate (10ml). The layers were (0.22ml). After stirring for 1.5h at 20°C the mixture was partitioned between 2M hydrochloric combined organic extracts were washed with water (20ml) and brine (20ml), dried over sodium sulphate and evaporated in vacuo. The crude material was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol (10:1) to give a pale yellow solid. To a solution of this in methanol (0.5ml) was added 1M sodium hydroxide acid (5ml) and dichloromethane (10ml). The layers were separated and the aqueous phase was further extracted with dichloromethane (2 \times 10ml). The combined organic extracts were vacuo to give the title compound as a pale yellow foam (0.123g, 73%). LCMS: R, 3.84 min; washed with water (20ml) and brine (20ml), dried over sodium sulphate and evaporated in

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Example 2: (2S)-2-(((2S)-4-Methyl-2-[(2-[(3-(1-piperidinylcarbonyl)-2-naphthyl] oxy}acetyl)amino]pentanoyl}amino)-3-{4-[({4-[(2-phenylacetyl)amino}-1piperidinyl)carbonyl)oxy]phenyl)propanoic acid

atmosphere, was added a solution of Intermediate 3 (0.2g) in anhydrous THF (2ml) followed To a solution of triphosgene (0.04g) in anhydrous dichloromethane (1ml), under a nitrogen

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by diisopropylethylamine (0.07ml). After stirring for 3h at 20°C Intermediate 59 (0.09g) was added followed by dilsopropylethylamine (0.07ml). Stirring was continued for 18h then the mixture was partitioned between 2M hydrochloric acid (30ml) and ethyl acetate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (20ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (20ml), water (20ml) and brine (20ml), dried over sodium sulphate and evaporated in vacuo. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate switching to ethyl acetate/ethanol (9:1) to give a white foam (0.19g). To a solution of this (0.15g) in methanol (2ml) was added 2M sodium hydroxide (0.18ml). After stirring for 1h at 20°C the mixture was partitioned between 2M hydrochloric acid (40ml) and ethyl acetate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (30ml). The comblned organic extracts were dried over sodium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:1) to give the title compound as a white solid (0.12g, 54% from Intermediate 3). LCMS: R, 3.73 min; m/z 834 (MH*).

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Example 3: (2S)-3-{4-[((4-[(2,2-Dicyclohexylacetyl)amino]-1-piperidinyl)carbonyl) oxy]phenyf}-2-{[(2S)-4-methyt-2-({2-[4-(1-piperidin; |carbonyl)phenoxy]acetyl}

amino)pentanoyijamino)propanoic acid

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(0.04g) and 1-hydroxybenzotriazole (0.03g). After stirring for 30 mins at 20°C Intermediate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl in methanol (2ml) was added 2M sodium hydroxlde (0.18ml). After stirring for 1h at 20°C the To a solution of Intermediate 48 (0.05g) in anhydrous DMF (3ml), under a nitrogen added 1-(3-dimethylaminopropyl)-3-ethylcarbodilmide hydrochloride 10 (0.13g) was added followed by diisopropylethylamine (0.08ml), and stirring was continued for 18h. The mixture was partitioned between 2M hydrochloric acid (40ml) and ethyl acetate acetate (30ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (30ml), water (2 x 30ml) and brine (20ml), dried over sodium sulphate mixture was partitioned between 2M hydrochloric acld (40ml) and ethyl acetate (30ml). The and evaporated in vacuo to give a cream coloured solid (0.16g). To a solution of this (0.15g) ayers were separated and the aqueous phase was further extracted with ethyl acetate vacuo. The crude product was purified by flash column chromatography on silica gel eluting (30ml). The combined organic extracts were dried over sodium sulphate and evaporated in Was

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with chloroform/methanol/acetic acid (95:5:1) to give the title compound as (0.12g, 62% from Intermediate 10). LCMS: R, 4.26 min; m/z 872 (MH*).

Example 4: (2S)-2-((2S)-4-Methyl-2-((2-[4-(1-piperidInylcarbonyl)phenoxy)

To a solution of Intermediate 48 (0.06g) in acetonitrile (5ml), under a nitrogen atmosphere, hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate 15 (0.1g) was added, and stirring was continued for 18h. The mixture was partitioned between water (20ml) dichloromethane/ethanol/880 ammonia (250:8:1) to give a white sticky solid (0:1g). To this material was purified by flash column chromatography on silica gel eluting with was added trifluoroacetic acid (3ml) and water (3 drops). After stirring for 4h at 20°C the and ethyl acetate (25ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (20ml). The combined organic extracts were washed with water (20ml) and brine (20ml), dried over sodium sulphate and evaporated in vacuo. The crude solvent was evaporated in vacuo and the residue was triturated with ether to give the title was added 1-(3-dimethylaminopropyl)-3-ethylcarbodilmide hydrochloride (0.06g) and acetyl}amino)pentanoyl]amino}-3-{4-{(4-morpholinylcarbonyl)oxy]phenyl} propanoic acid compound as a white solid (0.06g, 50%). LCMS: R, 3.21 min; m/z 653 (MH*) S

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methyl-2-((2-[4-(1-piperidinylcarbonyl)phenoxy]acetyl]amino)pentanoyl] amino)propanoic Example 5: (2S)-3-[4-([i4-(Aminocarbonyl)-1-piperidinyl]carbonyl)oxy)phenyl]-2-[[(2S)-4-

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crude Intermediate ester was purified by flash column chromatography on silica gel eluting with dichloromethane/ethanol/880 ammonia (500:8:1 switching via 250:8:1 to 100:8:1). The This was similarly prepared from Intermediate 48 (0.06g) and Intermediate 16 (0.12g). title compound was obtained as a white solid (0.09g, 59%). LCMS: R, 2.84 min; m/z

Example 6: (2S)-3-[4-([[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-[((2S)-2-[[2

This was similarly prepared from Intermediate 49 (0.07g) and Intermediate 16 (0.11g). The crude intermediate ester was purified by flash column chromatography on silica gel eluting with dichloromethane/ethanol/880 ammonia (500:8:1 switching via 250:8:1 to 100:8:1). The title compound was obtained as a white solid (0.08g, 42%). LCMS: R, 3.16 min; m/z (2-benzoylphenoxy)acetyl]amino}-4-methylpentanoyl)amino] propanoic acid

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Example 7: (2S)-2-[[(2S)-2-({2-[4-(Aminocarbonyl)phenoxy]acetyl}amino)-4-

methylpentanoyi]amino}-3-[4-([[4-(aminocarbonyl]-1-piperidinyl]carbonyl]oxy)

enviloropanole acid

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This was similarly prepared from intermediate 51 (0.06g) and intermediate 16 (0.11g). The crude intermediate ester was purified by flash column chromatography on silica gel eluting with dichloromethane/ethanol/880 ammonia (500:8:1 switching via 250:8:1 and 100:8:1 to 75:8:1). The title compound was obtained as a white solid (0.07g, 55%). LCMS: R, 2.65 min;

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Example 8: (2S)-3-{4-[((4-[(2-Cyclohexylacetyl)amino]-1-piperidinyl}carbonyl) oxylphenyl)-2-[((2S)-2-[(2-(2-iodophenoxy)acetyl]amino)-4-methylpentanoyl) amino]propanoic acid

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To a solution of triphosgene (0.058g) in anhydrous dichloromethane (2mi), under a nitrogen atmosphere, was added a solution of Intermediate 4 (0.246g) in anhydrous THF (2mi) followed by diisopropylethylamine (0.11ml). After stirring for 4h at 20°C (Intermediate 57 (0.1g) was added followed by diisopropylethylamine (0.07ml). Stirring was continued for 18h then the mixture was partitioned between 2M hydrochloric acid (50ml) and dichloromethane (50ml). The layers were separated and the organic extract was washed with water (20ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (1:1) to give a white foam (0.13g). To a solution of this (0.12g) in methanol (3ml) was added 2M sodlum hydroxide (1ml) and water (2ml). After stirring for 18h at 20°C the mixture was partitioned between 2M hydrochloric acid (30ml) and chloroform (30ml). The layers were separated and the organic phase was washed with water (20ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash, column chromatography on silica gel eluting with chloroform/methanol (4:1) to give the title compound as a white solid (0.064g, 20%). LCMS: R, 4.12 min; *m*Z 805 (MH*).

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Example 9: (2S)-3-{4-[((4-[(2,2-Dicyclohexylacetyl)amino]-1-piperidinyl)carbonyl) oxy]phenyl)-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methylpentanoyl)

aminolpropanoic acid

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This was similarly prepared from Intermediate 4 (0.203g) and Intermediate 58 (0.14g). The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol (9:1) to give the title compound as a white foam (0.153g, 52%). LCMS: R, 4.45 min; m/z 887 (MH*).

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Example 10: (2S)-2-(((2S)-2-((Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl)amino)-3-(4-[(4-morpholinylcarbonyl)oxy]phenyl)propanoic acid

To a solution of Intermediate 6 (0.165g) in dichloromethane (5ml), under a nitrogen atmosphere, was added morpholine (0.04ml) and diisopropylethylamine (0.05ml). After stirring for 30 mins at 20°C the solution was diluted with dichloromethane (50ml) and washed with saturated aqueous potassium carbonate (3 x 30ml), 1M hydrochloric acid (2 x 40ml) and water (30ml), dried over magnesium sulphate and evaporated in vacuo to give a white foam (0.143g). To a solution of this (0.14g) in methanol (2ml) was added 1M sodium hydroxide (2ml) and the mixture was stirred for 30 mins at 20°C, then partitioned between 1M hydrochloric acid (40ml) and ethyl acetate (50ml). The organic extract was washed with brine (30ml), dried over magnesium sulphate and evaporated in vacuo. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol (4:1) to give the title compound as a white solid (0.1g, 69%). LCMS: R, 3.85 min; m/z 602 (MH*).

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Example 11: (2S)-2-(((2S)-2-[(Dibenzolb,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl}amino)-3-[4-(((4-(2-furoyl)-1-piperazinyl)carbonyl)oxy)phenyl] propanoic acid To a solution of Intermediate 6 (0.13g) in dichloromethane (5ml), under a nitrogen atmosphere, was added 1-(2-furoyl)piperazine (0.04g) and diisopropylethylamine (0.04ml). After stirring for 3h at 20°C the solution was diluted with dichloromethane (20ml) and washed with saturated aqueous potassium carbonate (3 x 20ml), 1M hydrochloric acid (2 x 20ml) and water (20ml), dried over magnesium sulphate and evaporated *in vacuo* to give a white foam (0.153g). To a solution of this (0.15g) in methanol (2ml) was added 1M sodium hydroxide (2ml) and the mixture was stirred for 30 mlns at 20°C, then partitioned between 1M hydrochloric acid (20ml) and ethyl acetate (20ml). The organic extract was washed with brine (20ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol (4:1) to give the title compound as a white solid (0.126g, 92%). LCMS: R, 3.85 min; *m*/z 695 (MH+).

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Example 12: (2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-({(2S)-2-[(dibenzo[b,d]furan-4-yicarbonyl)amlno]-4-methylpentanoyl}amino)propanoic acid

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To a solution of Intermediate 6 (0.172g) in dichloromethane (4ml), under a nitrogen atmosphere, was added Intermediate 56 (0.084g) and diisopropylethylamine (0.2ml). After stirring for 3h at 20°C the solution was diluted with dichloromethane (50ml) and washed with

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water (50ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (4:1) to give a white foam. To a solution of this in methanol (2ml) was added 1M sodium hydroxide (2ml) and the mixture was stirred for 1h at 20°C, then partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (50ml). The organic extract was washed with brine (50ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol (4:1) to give the title compound as a white solid (0.041g, 23%), LCMS: R, 3.72 mln; *mz* 705 (MH*).

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Example 13: (2S)-2-(((2S)-2-((Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl)amino)-3-(4-[((4-[(2-phenylacetyl)amino]-1-piperidinyl)carbonyl)oxy] phenyl)propanolc acid

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To a solution of Intermediate 45 (0.055g) in acetonitrile (2ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.052g) and 1-hydroxybenzotriazole (0.038g). After stirring for 30 mins at 20°C Intermediate 8 (0.15g) was added followed by dilsopropylethylamine (0.047ml), and stirring was continued for 18h. The mixture was diluted with chloroform (100ml) and washed with 1M hydrochloric acid (3 x 50ml), saturated aqueous sodium hydrogen carbonate (3 x 50ml) and water (50ml), dried over magnesium sulphate and evaporated *in vacuo* to give a white foam (0.189g). To a solution of this (0.176g) in methanol (4ml) was added 1M sodium hydrochloric acid (50ml) and ethyl acetate (200ml). The organic extract was washed with brine (30ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with a gradient of chloroform/methanol (4:1) to give the title compound as a white solid (0.103g, 79%). LCMS: R, 4.00 min; *m*/z 733 (MH*).

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Example 14: (2S)-2-[((2S)-2-[(2-lodophenoxy)acetyl]amino]-4-methyl pentanoyi)amino]-3-[4-[(4-[(2-phenylacetyl)amino]-1-piperidinyl)carbonyl)oxyl phenyl)propanoic acid
This was similarly prepared from Intermediate 43 (0.073g) and Intermediate 8 (0.15g). The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol (6:1) to give the title compound as a white solid (0.103g, 53%). LCMS: R, 3.84 min; m/z 799 (MH*).

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Example 15: (2S)-3-(4-[[(4-Acetyl-1-piperazinyl)carbonyl]oxy]phenyl)-2-[((2S)-2-[[2-(2-iodophenoxy)acetyl]amino]-4-methylpentanoyl)amino]propanolc acid

To a solution of Intermediate 43 (0.07g) in acetonitrile (5ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.05g) and 1-hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate 21 (0.135g) was added followed by diisopropylethylamine (0.05mi) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (30ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (2 × 50ml), dried over sodium sulphate and evaporated in vacuo. The residue was co-evaporated with dichloromethane to give a white foam. To this was added trifluoroacetic acid (2ml) and water (3 drops). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white solid (0.143g, 83%). LCMS: R, 3.12 min; m/z 709 (MH*).

Example 16: (2S)-3-(4-[[(4-Acetyl-1-piperazinyl)carbonyl]oxy)phenyl)-2-[[(2S)-2-([2-[2-(tert-butyl)phenoxy]acetyl]amino)-4-methylpentanoyl]amino)propanoic acid

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To a solution of Intermediate 46 (0.052g) in acetonitrile (5ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodilmide hydrochloride (0.05g) and 1-hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate 21 (0.135g) was added followed by diisopropylethylamine (0.05ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate (30ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (2 × 50ml), dried over sodium sulphate and evaporated in vacuo. The residue was co-evaporated with dichloromethane to give a white foam. To this was added trifluoroacetic acid (2ml) and water (3 drops). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white solid (0.115g, 74%). LCMS: R, 3.31 min; mz 639 (MH*).

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Example 17: (2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-[((2S)-4-methyl-2-{[2-(2-methylphenoxy)acetyl]amino}pentanoyl)amino]propanoic acid

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To a solution of (2-methylphenoxy)acetic acid (0.042g) in acetonitrile (5ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.05g) and 1-hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate

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21 (0.135g) was added followed by diisopropylethylamine (0.05ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50mt) and ethyl acetate (30ml). The tayers were separated and the aqueous phase was further extracted with ethyl acetate (30ml). The combined organic extracts were washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (2 imes 50ml), dried over sodium sulphate and evaporated in vacuo. The residue was co-evaporated with dichloromethane to give a white foam. To this was added trifluoroacetic acid (2mi) and water (3 drops). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue triturated with ether to give the title compound as a white solid (0.124g, 86%). LCMS: R, 3.10

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Example 18: (2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-({(2S)-2-

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[(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl)amino)propanoic acid

To a solution of Intermediate 45 (0.053g) in acetonitrile (5ml), under a nitrogen atmosphere, hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate 21 (0.135g) was added followed by diisopropylethylamine (0.05ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (30ml). The was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.05g) and 1layers were separated and the aqueous phase was further extracted with ethyl acetate (30ml). The combined organic extracts were washed with saturated aqueous sodium evaporated in vacuo. The residue was co-evaporated with dichloromethane to give a white foam. To this was added trifluoroacetic acid (2ml) and water (3 drops). After stirring for 4h at hydrogen carbonate (40ml) and water (2 \times 50ml), dried over sodium sulphate and 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white solid (0.127g, 83%). LCMS: R, 3.33 min; m/z 643 (MH*).

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Example 19: (2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{((2S)-2-{[2-(2lodophenoxy)acetyl]amino}-4-methylpentanoyi)amino]propanoic acid

This was similarly prepared from Intermediate 43 (0.07g) and Intermediate 22 (0.151g). The title compound was obtained as a white solid (0.152g, 81%). 30

LCMS: R, 3.58 min; m/z 771 (MH*).

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Example 20: (2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{[(2S)-2-((2-{2 (tert-butyl)phenoxy]acetyl}amino)-4-methylpentanoyl]amino)propanoic acid

hydroxybenzotriazole (0.04g). After stirring for 30 mins at 20°C Intermediate 22 (0.151g) was To a solution of Intermediate 46 (0.052g) in acetonitrile (5ml), under a nitrogen atmosphere, added followed by diisopropylethylamine (0.05ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (30ml). The layers were separated and the aqueous phase was further extracted with ethyl acetate foam. To this was added trifluoroacetic acid (2ml) and water (3 drops). After stirring for 4h at hydrogen carbonate (40ml) and water (2 x 50ml), dried over sodium sulphate and evaporated in vacuo. The residue was co-evaporated with dichloromethane to give a white 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white foam (0.17g, 90%). LCMS: R, 3.61 min; m/z 701 (MH*). was added 1-(3-dimethylaminopropyl)-3-ethylcarbodilmide hydrochloride (0.05g) (30ml). The combined organic extracts were washed with saturated aqueous

Example 21: (2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{((2S)-4-methyl-2-[[2-(2-methylphenoxy)acetyl]amino)pentanoyl)amino]propanoic acid

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solution of (2-methylphenoxy)acetic acid (0.472g) in acetonttrile (30ml), under a hydrochloride (0.56g) and 1-hydroxybenzotriazole (0.4g). After stirring for 30 mins at 20°C a 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide solution of Intermediate 22 (1.5g) in acetonitrile (25ml) was added and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (75ml). The layers were separated and the organic phase was washed with saturated and evaporated in vacuo to give a white foam. To a solution of this in chloroform (12ml) was aqueous sodium hydrogen carbonate (40ml) and water (50ml), dried over sodium sulphate added trifluoroacetic acld (6ml). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was co-evaporated with chloroform and ether to compound as a white foam (0.17g, 90%). LCMS: R, 3.44 min; m/z 659 (MH*), added was atmosphere,

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Example 22: (2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)\Larbonyl]oxy)phenyl)-2-{((2S)-2-{[2-(2,4dichlorophenoxy)acetyljamino}-4-methyfpentanoyl)amino]propanoic acid 8

This was similarly prepared from 2,4-dichlorophenoxyacetic acid (0.055g) and Intermediate 22 (0.151g). The title compound was obtained by trituration with ether as a white solid (0.129g, 75%). LCMS: R, 3.52 min; m/z 713 (MH*).

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Example 23: (2S)-2-[((2S)-2-[[2-(2-lodophenoxy)acetyl]amino)-4-methyl pentanoyl)amino]-3-[4-[(4-morpholiny/carbonyl)oxy]phenyl}propanoic acid

To a solution of Intermediate 43 (0.556g) in acetonitrile (40ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.383g) and 1-hydroxybenzotriazole (0.27g). After stirring for 30 mins at 20°C Intermediate 23 (1g) was added followed by diisopropylethylamine (0.35ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (75ml). The layers were separated and the organic phase was washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (50ml), dried over sodium sulphate and evaporated in vacuo to give a white foam. To a solution of this in dichloromethane (20ml) was added trifluoroacetic acid (20ml) and water (1ml). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white solid (1.15g, 92%). LCMS: R, 3.68 min; m/z 668 (MH*).

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Example 24: (2S)-2-{[(2S)-2-({2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino)-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acid

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To a solution of intermediate 46 (0.416g) in acetonitrile (40ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.383g) and 1-hydroxybenzotriazole (0.27g). After stirring for 30 mins at 20°C intermediate 23 (1g) was added followed by dilsopropylethylamine (0.35ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (75ml). The layers were separated and the organic phase was washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (50ml), dried over sodium sulphate and evaporated in vacuo to give a white foam. To a solution of this in dichloromethane (20ml) was added trifluoroacetic acid (20ml) and water (1ml). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was triturated with ether to give the title compound as a white solid (0.63g, 53%). LCMS: R, 3.90 min; m/z 598 (MH+).

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NMR (DMSO-d₆) 5H 12.74 (br s, 1H), 8.38 (d, 1H), 7.81 (d, 1H), 7.20-7.25 (m's, 3H), 7.14 (m, 1H), 6.99 (d, 2H), 6.90 (m, 1H), 6.85 (d, 1H), 4.57 (d, 1H), 4.50 (m's, 3H), 3.61 (m, 4H), 3.52 (br m, 2H), 3.30-3.40 (excess 2H, obscured by water), 3.06 (dd, 1H), 2.90 (dd, 1H), 1.57 (m, 1H), 1.38-1.50 (m's, 2H), 1.35 (s, 9H), 0.87 (d, 3H), 0.85 (d, 3H).

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Example 24 (Alternative Procedure): (2S)-2-((212-(Tert-butyl)phenoxylacetyl) amino)-4-methyl pentanoyl]amino)-3-(4-((4-morpholinylcarbonyl)oxylphenyl)propanoic acid To Sasrin resin (125g) was added a solution of (2S)-3-(4-(allyloxy)phenyl)-2-((9H-fluoren-9-ylmethoxy)carbonyl]amino)propanoic acid (300g) in DMF (970ml). After 15 mins pyridine (60ml) was added followed by 2,6-dichlorobenzoyl chloride (106.5ml) dropwise. The mixture was stirred for 18h at 20°C. The resin was filtered and washed with DMF (3 x 800ml), methanol (3 x 800ml) and dichloromethane (3 x 11). The resin was treated with acetic anhydride (800ml) and pyridine (10ml) and the mixture was stirred for 3.5h at 45°C. After cooling to 20°C the resin was filtered and washed with NMP (3 x 800ml), methanol (3 x 800ml) and dichloromethane (3 x 800ml) then dried *in vacuo*.

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200g of the resin was treated with 20% piperidine in DMF (1.2l) and stirred for 3h at 20°C. The resin was filtered and washed with DMF (3 x 1l), methanol (3 x 1l) and dichloromethane (3 x 1l). To this was added a solution of Fmoc-leucine (233.3g), 1,3-dilsopropylcarbodiimide (84.7g) and 1-hydroxybenzotriazole (89.3g) in NMP (1.2l). The mixture was stirred for 18h at 20°C. The resin was filtered and washed with NMP (3 x 1l), methanol (3 x 1l) and dichloromethane (3 x 1l).

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The resin was treated with 20% piperidine in DMF (1.2l) and stirred for 3h at 20°C. The resin was filtered and washed with DMF (3 x 1l), methanol (3 x 1l) and dichloromethane (3 x 1l). To this was added a solution of Intermediate 46 (68.8g), 1,3-diisopropylcarbodiimide (42.3g) and 1-hydroxybenzotriazole (44.7g) in NMP (1.2i). The mixture was stirred for 18h at 20°C. The resin was filtered and washed with NMP (3 x 1l), methanol (3 x 1l) and dichloromethane

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To the resin was added dichloromethane (500ml), phenylsilane (160ml) and a slurry of tetrakis(triphenylphosphine)palladium(0) (34g) in dichloromethane (500ml). The mixture was stirred for 2h at 20°C. The resin was filtered and washed with dichloromethane (3 x 1l), ether (3 x 1l) and dichloromethane (6 x 1l).

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A slurry of the resin in dichloromethane (800mi) was treated with dilsopropylethylamine (120ml) followed by 4-nitrophenyl chloroformate (131g) in 3 portions at 10 minute intervals. The mixture was stirred for 2h at 20°C. The resin was filtered and washed with dichloromethane (3 x 1l), ether (3 x 1l) and DMF (3 x 1l). A slurry of the resin in DMF (800ml) was treated with a solution of morpholine (56.5ml) in DMF (200ml). The mixture was stirred for 2h at 20°C. The resin was filtered and washed with DMF (3 x 1l), ether (3 x 1l) and dichloromethane (3 x 1l).

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A slurry of the resin in dichloromethane (400ml) was treated with 10% TFA in dichloromethane (800ml). After stirring for 30 mins at 20°C the resin was filtered and washed

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with dichloromethane (2 x 500 ml). The combined filtrate and washings were evaporated in vacuo. The residue was triturated with ether (750ml) and the resulting white solid filtered. To this was added acetonitrile (500ml) and the mixture was heated to reflux. The hot solution was filtered and the filtrate allowed to cool to 20°C. The mixture was filtered to give the title compound as a white solid (50.9g).

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Example 25: (2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino]-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acld

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To a solution of (2-methylphenoxy)acetic acid (0.332g) in acetonitrile (40ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.383g) and 1-hydroxybenzotriazole (0.27g). After stirring for 30 mins at 20°C Intermediate 23 (1g) was added followed by dilsopropylethylamine (0.35ml) and stirring was continued for 18h. The mixture was partitioned between 1M hydrochloric acid (50ml) and ethyl acetate (75ml). The layers were separated and the organic phase was washed with saturated aqueous sodium hydrogen carbonate (40ml) and water (50ml), dried over sodium sulphate and evaporated *in vacuo* to give a white foam. To a solution of this in dichloromethane (20ml) was added trifluoroacetic acid (20ml) and water (1ml). After stirring for 4h at 20°C the solvent was evaporated *in vacuo* and the residue was triturated with ether to give the title compound as a white solid (0.895g, 80%). LCMS: R₁ 3.31 min; *m/z* 556 (MH*).

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Example 28: (2S)-3-[4-([[4-(AmInocarbonyl)-1-piperidinyl]carbonyl)oxy)phenyl]-2-[((2S)-2-([2-(2-lodophenoxy)acetyl]amino)-4-methylpentanoyl)amino]propanoic acid

This was similarly prepared from intermediate 43 (0.06g) and intermediate 24 (0.1g). The title compound was obtained as a white solid (0.07g, 56%).

LCMS: R, 3.33 min; m/z 709 (MH*).

Example 27: (2S)-3-[4-({[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-[((2S)-4-methyl-2-{[2-(2-methylphenoxy)acetyl]amino}pentanoyl)amino] propanoic acid

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To a solution of (2-methylphenoxy)acetic acid (0.345g) in acetonltrile (50ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.4g) and 1-hydroxybenzotriazole (0.3g). After stirring for 30 mins at 20°C Intermediate 24 (1g) was added followed by diisopropylethylamine (0.35ml) and stirring was continued for 18h. The mixture was concentrated *in vacuo* and the residue was partitioned between 1M hydrochloric acid (100ml) and ethyl acetate (300ml). The layers were separated

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and the organic phase was washed with 1M hydrochloric acid (2 × 100ml), saturated aqueous sodium hydrogen carbonate (3 × 100ml) and brine (100ml), dried over magnesium sulphate and evaporated *in vacuo* to give a white solid. To a solution of this in chloroform (5ml) was added trifluoroacetic acid (5ml) and water (1ml). After stirring for 3h at 20°C the solvent was evaporated *in vacuo* and the residue was azeotroped with toluene (2 × 20ml) then triturated with ether to give the title compound as a white solid (1.05g, 96%). LCMS: R₁ 3.20 min; *m/z* 597 (MH*). Solubility in water. 0.01 mg/ml.

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NMR (DMSO-d₆) 5H 12.75 (br s, 1H), 8.33 (d, 1H), 7.81 (d, 1H), 7.32 (br s, 1H), 7.21 (d, 2H), 7.15 (d, 1H), 7.11 (t, 1H), 6.98 (d, 2H), 6.79-6.89 (m's, 3H), 4.46-4.56 (AB system, 2H), 4.39-4.46 (m's, 2H), 3.95-4.14 (m's, 2H), 2.80-3.10 (m's, 4H), 2.33 (m, 1H), 2.20 (s, 3H), 1.75 (m, 2H), 1.40-1.60 (m's, 5H), 0.82-0.87 (m's, 6H).

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Example 27 (Alternative Procedure): (2S)-3-[4-(([4-(Aminocarbonyl)-1-piperidinyl]carbonyl)oxy)phenyl]-2-[((2S)-4-methyl-2-[(2-methylphenoxy)acetyl]amino)pentanoyl)amino)

To Wang resin (50g) was added a solution of (2S)-3-[4-(allyloxy)phenyl]-2-[(terf-butoxycarbonyl)amino]propanoic acid (115.8g) and 1-hydroxybenzotriazole (48.6g) in DMF (475ml). After 15 minutes 1,3-diisopropylcarbodiimide (56.5ml) was added and the mixture was stirred for 24h at 45°C. The resin was filtered and washed with DMF (3 x 360ml), methanol (3 x 360ml) and dichloromethane (3 x 700ml). To a slurry of the resin in dichloromethane (644ml) was added pyridine (14.7ml). Acetic anhydride (26.9ml) was added and the mixture was stirred for 12h at 20°C. The resin was filtered and washed with dichloromethane (3 x 550ml), methanol (3 x 370ml) and dichloromethane (3 x 550ml).

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A slurry of 20g of the resin in dichloromethane (100ml) was cooled to 2-5°C and treated with a solution of phenol (20g) in dichloromethane (80ml). Chlorotrimethylsilane (20ml) was added dropwise and the mixture was stirred for 6h at 2-5°C. The resin was filtered and washed with dichloromethane (3 x 200ml), methanol (3 x 200ml), 10% water in DMF (2 x 200ml), 10% diisopropylethylamine in DMF (3 x 200ml), DMF (200ml), methanol (3 x 200ml).

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A slurry of the resin in DMF (55ml) was treated with a solution of Fmoc-leucine (32.7g) and 1-hydroxybenzotriazole (12.5g) in DMF (85ml). After 5 minutes 1,3-diisopropylcarbodiimide (19.3ml) was added and the mixture was stirred for 15h at 20°C. The resin was filtered and washed with DMF (3 x 150ml), methanol (3 x 150ml) and dichloromethane (3 x 150ml).

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The resin was treated with 20% piperidine in DMF (180ml) and stirred for 1h at 20°C. The resin was filtered and washed with DMF (3 x 150ml), dichloromethane (3 x 150ml), DMF (3 x

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150ml) and dichloromethane (3 x 150ml). To a sturry of this in DMF (50ml) was added a solution of (2-methylphenoxy)acetic acid (17.9g) and 1-hydroxybenzotriazole (14.6g) in DMF (100ml). After 5 minutes 1,3-diisopropylcarbodilmide (16.9ml) was added and the mixture was stirred for 65h at 20°C. The resin was filtered and washed with DMF (2 x 150ml), methanol (3 x 150ml) and dichloromethane (3 x 150ml).

A slurry of the resin in dichloromethane (60ml) was treated with a solution of tetrakls(triphenylphosphine)palladium(0) (5.21g) in dichloromethane (140ml) followed by morpholine (13ml). The mixture was stirred for 2h at 20°C then the resin was filtered and washed with dichloromethane (7 x 200ml).

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A slurry of the resin in dichloromethane (160ml) was treated with dilsopropylethylamine (12.4ml) followed by 4-nitrophenyl chloroformate (24.8g) in 3 portions at 5 minute intervals. The mixture was stirred for 1h at 20°C. The resin was filtered and washed with dichloromethane (3 x 200ml). The resin was treated with a solution of isonipecotamide (15.8g) in DMF (180ml) and the mixture was stirred for 1.5h at 20°C. The resin was filtered and washed with DMF (4 x 200ml) and dichloromethane (2 x 200ml).

The resin was treated with 50% TFA In dichloromethane (200ml). After stirring for 1h at 20°C the resin was filtered and washed with dichloromethane (5 x 200 ml). The combined filtrate and washings were evaporated *in vacuo*. The residue was azeotroped with toluene (2 x 100ml) then triturated with ether (50ml) and the resulting white solid filtered. To this was added acetonitrile (150ml) and the mixture was heated to reflux. The resulting suspension was allowed to cool to 20°C and stirred for 18h.. The mixture was filtered to give the title compound as a white solid (4.9g).

Example 27A: (2S)-3-[4-([[4-(Aminocarbonyl)-1-piperidinyl]carbonyl]oxy)phenyl]-2-[((2S)-4-methyl-2-([2-(2-methylphenoxy)acetyl]amino)pentanoyl)amino] propanolc acid potassium salt A suspension of Example 27 (10g) in methanol (150ml) was warmed to reflux to obtain a clear solution. To this was added a solution of potassium carbonate (1.16g) in water (7.5ml). After heating under reflux for two minutes the solvents were evaporated in vacuo to give a crisp foam. To this was added acetonitrile (100ml) and the mixture was warmed to reflux, during which time the foam collapsed and started to crystallise. After ten minutes the mixture was allowed to cool to 20°C then filtered under reduced pressure, washed with acetonitrile (25ml) and ether (50ml) to give the title compound as a white solid (10.65g, 100%). The product is believed to be isolated in the form of its monohydrate. Solubility in water: >250

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NMR (DMSO-d₆) 5H 8.27 (d, 1H), 7.42 (d, 1H), 7.37 (d, 1H), 7.04-7.16 (m's, 4H), 6.78-6.88 (m's, 5H), 4.44-4.59 (AB system, 2H), 4.21 (m, 1H), 3.95-4.12 (br m's, 2H), 3.87 (m, 1H), 2.80-3.10 (m's, 4H), 2.34 (m, 1H), 2.20 (s, 3H), 1.75 (m, 2H), 1.41-1.60 (m's 5H), 0.86 (d, 3H), 0.80 (d, 3H)

Example 28: (2S)-3-[4-([[4-(Aminocarbony!)-1-piperidinyl]carbonyl)oxy)phenyl]-2-(((2S)-2-[(dibenzo[b,d]furan-4-yicarbonyl)amino]-4-methylpentanoyl]amino) propanoic acid

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To a solution of Intermediate 45 (0.438g) in acetonitrile (50ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodifmide hydrochloride (0.4g) and 1-hydroxybenzotriazole (0.29g). After stirring for 30 mins at 20°C Intermediate 24 (1g) was added followed by dilsopropylethylamine (0.35ml) and stirring was continued for 18h. The mixture was concentrated *in vacuo* and the residue was partitioned between 1M hydrochloric acid (100ml) and ethyl acetate (300ml). The layers were separated and the organic phase was washed with 1M hydrochloric acid (2 × 100ml), saturated aqueous sodium hydrogen carbonate (3 × 100ml) and brine (100ml), dried over magnesium sulphate and evaporated *in vacuo* to give a white solid. To a solution of this in chloroform (5ml) was added trifluoroacetic acid (5ml) and water (1ml). After stirring for 3h at 20°C the solvent was evaporated *in vacuo* and the residue was azeotroped with toluene (2 × 20ml) then triturated with ether to give the title compound as a white solid (0.95g, 80%). LCMS: R, 3.48 min; *m/z* 643 (MH*).

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Example 29: (2S)-2-{[(2S)-2-((2-{2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl[amino]-3-{4-({4-(1-piperidinylcarbonyl)-1-piperidinyl]carbonyl]oxy) phenyl]propanoic acid

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To a solution of Intermediate 46 (0.1g) in acetonitrile (5ml), under a nitrogen atmosphere, was added 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (0.09g) and 1-hydroxybenzotriazole (0.063g). After stirring for 30 mlns at 20°C Intermediate 20 (0.18g) was added and stirring was continued for 18h. The mixture was partitioned between water (20ml) and ethyl acetate (20ml). The layers were separated and the organic phase was washed with saturated aqueous sodium hydrogen carbonate (2 × 30ml), water (30ml) and brine (30ml), dried over sodium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with dichloromethane (8ml) was added trifluoroacetic acld (2ml). After stirring for 2h at 20°C the solvent was evaporated *in vacuo* and the crude product purified by flash column chromatography on silica gel eluting with

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dichloromethane/methanol/acetic acid/water (240:15:3:2) to give the title compound as a white foam (0.08g, 36%). LCMS: R₁ 4.07 min; m/z 707 (MH*).

Example 30: (2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino] 3-[4-(([4-(1-piperidinylcarbonyl)-1-piperidinyl]carbonyl)oxy) phenyl]propanoic acld

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This was similarly prepared from (2-methylphenoxy)acetic acid (0.09g) and Intermediate 20 (0.3g). The crude product was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol/acetic acid/water (240:15:3:2) to give the title compound as a white foam (0.116g, 34%). LCMS: R, 3.56 min; m/z 665 (MH*).

Example 31: (2S)-2-(((2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl}amino)-3-[4-([[4-(1-piperidinylcarbonyl)-1-piperidinyl]carbonyl}oxy)

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This was similarly prepared from Intermediate 45 (0.1g) and Intermediate 20 (0.176g). The crude product was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol/acetic acid/water (180:15:3:2) to give the title compound as a white foam (0.075g, 35%). LCMS: R_r 4.09 min; *m/z* 711 (MH*).

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Example 32: (2S)-2-{((2S)-2-((2-[(1-Bromo-2-naphthyl)oxy]acetyl}amino)-4-methylpentanoyl]amino}-3-[4-(([4-(1-piperidinylcarbonyl)-1-piperidinyl]carbonyl)oxy)phenyl]propanoic acid

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This was similarly prepared from Intermediate 50 (0.124g) and Intermediate 20 (0.168g). The crude product was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol/acetic acid/water (200:15:3:2) to give the title compound as a white foam (0.055g, 24%). LCMS: R₄ 4.19 min; *m/z* 779 (MH*).

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Example 33: (2S)-3-[4-({[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-{[(2S)-2-((2-[2-(tert-butyl)phenoxy]acetyl)amino)-4-methylpentanoyl]amino} propanoic acid

To a solution of Intermediate 26 (0.47g) in dichloromethane (8ml), under a nitrogen atmosphere, was added isonipecotamide (0.106g) and disopropylethylamine (0.2ml). The mixture was stirred for 18h at 20°C then diluted with chloroform (100ml), washed with saturated aqueous potassium carbonate (3 x 50ml), 1M hydrochloric acid (3 x 50ml) and water (50ml), dried over magnesium sulphate and evaporated in vacuo to give a white foam. To a solution of this in chloroform (3ml) was added trifluoroacetic acid (3ml). After stirring for

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4h at 20°C the solvent was evaporated *in vacuo* and the residue was triturated with ether to give the title compound as a white solid (0.223g, 52%).

LCMS: R, 3.35 min; m/z 639 (MH*).

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Example 34: (2S)-2-{[(2S)-2-({2-{2-(Tert-butyl)phenoxyjacetyl}amino)-4-methyl pentanoyl]amino}-3-(4-{[(4-{[(4-{iuorobenzyl)amino]carbonyl}-1-piperidinyl) carbonyl]oxy}phenyl)propanoic acid

This was similarly prepared from Intermediate 26 (0.312g) and Intermediate 54 (0.181g). The title compound was obtained as a white solid (0.187g, 57%).

10 LCMS: R, 3.71 min; m/z 747 (MH*).

Example 35: (2S)-2-[((2S)-2-{[2-(2,4-Dichlorophenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acid

To a suspension of anhydrous potassium carbonate (0.057g) and sodium lodide (0.051g) in anhydrous DMF (1ml) was added 2,4-dichlorophenol (0.166g) followed by Intermediate 27 (0.2g). The mixture was stirred for 18h at 20°C then partitioned between saturated aqueous sodium hydrogen carbonate (10ml) and ethyl acetate (10ml). The layers were separated and the organic phase was further washed with saturated aqueous sodium hydrogen carbonate (10ml) and brine (10ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (1:1) to give a white foam. To a solution of this in dichloromethane (2ml) was added trifluoroacetic acid (2ml). After stirring for 2h at 20°C the solvent was evaporated *in vacuo* and the residue was triturated with ether to give the title compound as a white solid (0.146g, 70%). LCMS: R, 3.70 min; *mz* 610 (MH*).

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Example 36: (2S)-2-[((2S)-2-{[2-(2-Benzoylphenoxy)acetyl]amino}-4-methylpentanoyl)amino]-3-(4-[(4-morpholinylcarbonyl)oxy]phenyl}propanolc acid

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This was similarly prepared from 2-hydroxybenzophenone (0.2g) and Intermediate 27 (0.2g).

The title compound was obtained as a pale yellow foam (0.057g, 26%). LCMS: R, 3.60 min; m/z 646 (MH*).

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Example 37: (2S)-2-[((2S)-4-Methyl-2-{[2-(2-propylphenoxy)acetyt]amino} pentanoyl)amino}-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acid

This was similarly prepared from 2-propylphenol (0.14ml) and Intermediate 27 (0.2g). The title compound was obtained as a white solid (0.141g, 70%).

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LCMS: R, 3.71 mln; m/z 584 (MH1).

Example 38: (2S)-2-{[(2S)-2-((2-[(1-Bromo-2-naphthyl)oxy]acetyl}amino)-4-methylpentanoyl]amino}-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acid
This was similarly prepared from 1-bromo-2-naphthol (0.23g) and intermediate 27 (0.2g).
The title compound was obtained as a white solid (0.11g, 48%).
LCMS: R₁ 3.91 min; m/z 670 (MH*).

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Example 39: (2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl

pentanoyl)aminoj-3-(4-[(4-morpholinylcarbonyl)oxy]phenyl)propanoic acid

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To a suspension of anhydrous potassium carbonate (0.1g) and sodium iodide (0.06g) in anhydrous DMF (1ml) was added 2-cyclohexylphenol (0.12g) followed by Intermediate 27 (0.2g). The mixture was stirred for 18h at 20°C then partitioned between saturated aqueous sodium hydrogen carbonate (10ml) and ethyl acetate (10ml). The layers were separated and the organic phase was further washed with saturated aqueous sodium hydrogen carbonate (10ml) and brine (10ml), dried over magnesium sulphate and evaporated *in vacuo*. The crude material was purified by flash column chromatography on silica gel eluting with ethyl acetate/cyclohexane (1:1) to give a white foam. To a solution of this in dichloromethane (3ml) was added trifluoroacetic acid (3ml). After stirring for 2h at 20°C the solvent was evaporated *in vacuo* and the residue was azeotroped with toluene then triturated with ether to give the title compound as a white solid (0.118g, 55%). LCMS: R, 4.16 min; *m/z* 624 (MH*).

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Example 40: (2S)-2-[((2S)-2-{[(Benzyloxy)carbonyl]amino}-4-methylpentanoyl) amino]-3-{4- [(4-morpholinylcarbonyl)oxy]phenyl)propanoic acid

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To a solution of Intermediate 13 (0.19g) in chloroform (2ml) was added trifluoroacetic acid (2ml). After stirring for 4h at 20°C the solvent was evaporated in vacuo and the residue was trifurated with ether to give the title compound as a white solid (0.156g, 90%). LCMS: R, 3.22 min; m/z 542 (MH*).

Example 41: (2S)-3-[4-([[4-(2-Furoyl)-1-piperazinyl]carbonyl]oxy)phenyl]-2-[((2S) -2-([2-(2-lodophenoxy)acetyl]amino]-4-methylpentanoyl)amino]propanoic acid

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Intermediate 38 (0.26mmol) was treated with DMF (4ml). 2-lodophenol (0.57g), potassium carbonate (0.36g) and sodium iodide (0.39g) were added and the mixture was shaken for 16h at 20°C. The resin was filtered and washed with water (2 x 5ml), DMF (5 x 5ml) and

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dichloromethane (5 x 5ml) then treated with 1:1 trifluoroacetic acid/ dichloromethane (4ml). After 30 mins the resin was filtered and the filtrate was evaporated *in vacuo*. The residue was azeotroped with toluene (5ml) then triturated with ether. The crude product was crystallised from acetonitrile to give the title compound as a white solid (0.043g).

5 LCMS: R₁ 3.50 min; m/z 761 (MH⁺).

Example 42: (2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-[4-(([4-(2-furoyl)-1-piperazinyl]carbonyl]oxy)phenyl} propanoic acid

Intermediate 38 (0.26mmol) was treated with DMF (4ml). 2-tert-butyl phenol (0.4ml), potassium carbonate (0.36g) and sodium iodide (0.39g) were added and the mixture was shaken for 16h at 20°C. The resin was filtered and washed with vater (2 x 5ml), DMF (5 x 5ml) and dichloromethane (5 x 5ml) then treated with 1:1 trifluoroacetic acid/dichloromethane (4ml). After 30 mins the resin was filtered and the filtrate was evaporated in vacuo. The residue was azeotroped with toluene (5ml) then triturated with ether. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:1) to give the title compound as a white solid (0.04g). LCMS: R, 3.63 min; m/z 691 (MH*).

Example 43: (2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl

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pentanoyl)amlnoj-3-[4-([[4-(2-furoyl)-1-plperazinyl]carbonyl]oxy)phenyl] propanoic acid
This was similarly prepared from Intermediate 38 (0.26mmol) and 2-cyclohexyl phenol
(0.46g). The crude product was purified using a solid phase extraction cartridge containing
reverse phase silica eluting with a chloroform/methanol gradient (increasing from 98:2 to
80:20) to give the title compound as a cream solid (0.037g). LCMS: R, 3.83 min; m/z 717

25 (MH*).

Example 44: (2S)-2-{[(2S)-2-({2-|(1-Bromo-2-naphthyl)oxy]acetyl}amino)-4-methylpentanoyl]amino}-3-[4-([(4-(2-furoyl)-1-piperazinyl]carbonyl}oxy)phenyl] propanoic acid

This was similarly prepared from Intermediate 38 (0.26mmol) and 1-bromo-2-naphthol (0.58g). The crude product was crystallised from acetonitrile to give the title compound as a cream coloured solid (0.064g). LCMS: R, 3.69 min; m/z 763 (MH*).

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Example 45: (2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-plperidinyl) carbonyl]oxy}phenyl)-2-{((2S)-2-{[2-(2-cyclohexylphenoxy)acetyl]amino}-4-methylpentanoyl)amino]propanoic acid

This was similarly prepared from intermediate 39 (0.29mmol) and 2-cyclohexyl phenol (0.48g). The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:0.5) to give the title compound as a white solid (0.073g). LCMS: R, 4.13 min; m/z 789 (MH*).

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Example 46: (2S)-2-[((2S)-2-([2-(2-Benzoylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-(4-([(4-{[2-(4-chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy)phenyl)propanoic acid

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This was similarly prepared from intermediate 39 (0.29mmol) and 2-hydroxybenzophenone (0.55g). The crude product was purified by flash column chromatography on silica get eluting with chloroform/methanol/acetic acid (95:5:0.5) to give the title compound as a white solid (0.065g). LCMS: R, 3.75 min; m/z 811 (MH*).

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Example 47: (2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy)phenyl)-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methyl pentanoyl)amino]propanolc acid

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Intermediate 37 (0.27mmol) was treated with 20% piperidine in DMF (5ml) and shaken for 1h at 20°C. The resin was filtered and washed with DMF (5 x 5ml). A solution of Intermediate 43 (0.154g) in DMF (3ml) was added followed by a solution of benzotriazol-1-yl-oxy-trispyrrolidinophosphonium hexafluoro phosphate (0.285g) in DMF (2ml) and diisopropylethylamine (0.26ml). The mixture was shaken for 18h at 20°C. The resin was filtered and washed with DMF (5 x 5ml) and dichloromethane (5 x 5ml), then treated with 1:1 trifluoroacetic acid/ dichloromethane (5ml). After 30 mins the resin was filtered and the filtrate was evaporated *In vacuo*. The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:0.5) to give the title compound as a white solid (0.083g). LCMS: R₄ 3.76 min; *m/z* 833 (MH*).

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Example 48: (2S)-2-{[(2S)-2-({2-{2-(Tert-butyl)phenoxy}acetyl}amino}-4-methyl pentanoyl]amino}-3-(4-{[(4-{[2-(4-chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy}phenyl)propanolc acid

3

This was similarly prepared from intermediate 37 (0.27mmol) and intermediate 46 (0.115g). The crude product was purified by flash column chromatography on silica gel eluting with

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LCMS: R, 3.99 min; m/z 729 (MH*).

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chloroform/methanol/acetic acld (95:5:0.5) to give the title compound as a white solid (0.107g). LCMS: R, 3.93 min; m/z 763 (MH*).

Example 49: (2S)-3-(4-[[(4-[[2-(4-Chlorophenyi)acetyl]amino}-1-plperidinyl)

5 carbonyl]oxy)phenyl)-2-({(2S)-2-{(dibenzo[b,d]furan-4-y/carbonyl)amino]-4-methylpentanoyl]amino)propanoic acid

This was similarly prepared from Intermediate 37 (0.27mmol) and Intermediate 45 (0.117g).

The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:0.5) to give the title compound as a white solid

10 (0.056g). LCMS: R, 3.80 min; m/z 765 [M-H];.

Example 50: (2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidinyl)carbonyl]oxy}phenyl)-2-({(2S)-4-methyl-2-[(2-{[3-(1-piperidinylcarbonyl)-2-naphthyl]oxy}acetyl)amino]pentanoyl}amlno)propanoic acid

This was similarly prepared from Intermediate 37 (0.27mmol) and intermediate 44 (0.173g). The crude product was purified by flash column chromatography on silica gel eluting with chloroform/methanol/acetic acid (95:5:0.5) to give the title compound as a white solid (0.062g). LCMS: R, 3.71 min; m/z 868 (MH*).

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Example 51: (2S)-2-{[(2S)-2-({2-{2-(Tert-butyl)phenoxy}acetyl}amino}-4-methyl pentanoyl]amino}-3-{4-[({4-{(2-{12-(Tert-butyl)phenoxy}acetyl}amino}-1-piperidlnyl}carbonyl) oxyl phenyl}propanoic acid Intermediate 33 (0.23mmol) was treated with 1:1 dichloromethane/THF (3ml). Intermediate

59 (0.105g) was added followed by dilsopropylethylamine (0.16ml). After shaking for 18h at 20°C the resin was filtered, washed with dichloromethane (4 x 5ml) and ether (3 x 5ml) and then dried *in vacuo.* LCMS showed that some of the 4-nitrophenyl carbonate had been hydrolysed to the phenol so the resin was treated with 1:1 dichloromethane/THF (3ml), diisopropylethylamine (0.2ml) and 4-nitrophenyl chloroformate (0.23g). After shaking for 18h at 20°C the resin was filtered and washed with dichloromethane (4 x 5ml) then treated with 1:1 dichloromethane/THF (3ml), Intermediate 59 (0.07g) and diisopropylethylamine (0.12ml). After shaking for 18h at 20°C the resin was filtered and washed with 1:1 trifluoroacetic acid/dichloromethane (3ml). After 30 mins the resin was filtered and the filtrate was evaporated *in vacuo.* The residue was co-evaporated with dichloromethane followed by ether to give the title compound as an off-white solid (0.083g).

R7

pentanoy]]amino}-3-{4-[({4-[(2-cyclohexylacetyl)amino}-1-piperidinyl)carbonyl)

Example 52: (2S)-2-{((2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl

oxy]phenyi}propanoic acid

5 This was similarly prepared from Intermediate 33 (0.23mmol) and Intermediate 57 (0.106g).

The title compound was obtained as an off-white solid (0.073g).

LCMS: R, 4.27 min; m/z 735 (MH*),

Example 53: (2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl

pentanoyl]amino}-3-{4-[((4-[(2,2-dlcyclohexylacetyl)amino]-1-piperidinyl}

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carbonyl)oxy]phenyl)propanolc acid

This was similarly prepared from Intermediate 33 (0.25mmol) and Intermediate 58 (0.144g)

The title compound was obtained as an off-white solid (0.105g).

LCMS: R, 4.63 min; m/z 817 (MH*).

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Example 54: (2S)-2-[((2S)-4-Methyl-2-[[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino]-

3-{4-[((4-[(2-phenylacetyl)amino]-1-plperidinyl)carbonyl) oxy]phenyl}propanoic acid

This was similarly prepared from intermediate 34 (0.3mmol) and intermediate 59 (0.196g).

The crude product was purified by flash column chromatography on silica gel eluting with dichloromethane/methanol/acetic acid/water (240:15:3:2) to give the title compound as a

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pale yellow foam (0.091g). LCMS: R₁ 3.49 min; m/z 687 (MH*).

Example 55: (2S)-2-[((2S)-2-[[2-(2-Cyclohexylphenoxy)acetyl]amino]-4-methyl

pentanoyt)amino]-3-(4-[((4-[(2-phenylacetyl)amino]-1-piperidinyl)carbonyl)

oxy]phenyl]propanoic acid

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Intermediate 42 (0.27mmol) was treated with a solution of Intermediate 59 (0.178g) in 1:1 dichloromethane/THF (2ml) followed by disopropylethylamine (0.95ml). After shaking for 2h at 20°C the resin was filtered and washed with dichloromethane (5 x 5ml) then treated with 1:1 trifluoroacetic acid/dichloromethane (3ml). After 30 mins the resin was filtered and the filtrate was evaporated *in vacuo*. The residue was triturated with ether to give the title compound as an off-white solid (0.074g). LCMS: R, 4.04 min; *m/z* 755 (MH*).

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Example 56: (2S)-3-{4-[((4-[(2-Cyclohexylacetyl)amino]-1-piperidinyl)carbonyl) oxylphenyl}-

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2-[((2S)-2-{[2-(2-cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]propanoic acid This was similarly prepared from Intermediate 42 (0.27mmol) and Intermediate 57 (0.18g).

The title compound was obtained as an off-white solid (0.102g).

LCMS: R, 4.22 min; m/z 761 (MH*).

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Biological Data

The compounds of the Examples were tested in assay (1), the Jurkat adhesion assay, and the results obtained were as follows:

		7		7	1		7	7 -	7		,			_											
	*c	ဖ	4	4	4	4	4	4	4	2	6	10	8	4	4	8	80	8	5	5	9	6	4	10	10
	SEM*	0.18	0.24	0.12	0.08	0.03	0.00	0.03	0.15	0.10	0.05	0.12	90.0	0.11	0.07	0.04	0.10	0.09	0.38	0.06	0.03	0.07	0.17	0.02	. 0.08
d were as rollows:	plCso	7.88	8.03	7.38	7.78	8.11	8.25	8.58	7.37	7.58	8.08	8.08	7.96	7.59	7.78	8.57	8.49	8.59	8.43	8.12	7.83	8.41	7.85	8.35	8.22
the results obtained were as follows:	Example	-	2	က	4	5	9	7	Φ.	6	10.	11	12	13	14	15	16	17	18	19	20	21	22	23	24

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n	10	4	7	10	9	4	4	9	6	4	9	9	9	4	4	4	. 4	9	9	5	9	4	4	4	4	4	8	4	လ	, lu	4	9
SEM*	0.08	0.03	0.10	0.05	0.08	0.03	0.04	0.13	0.03	0.14	90.0	0.07	0.07	0.15	0.12	0.15	0.07	0.02	0.07	0.05	0.11	0.04	0.07	0.04	90.0	0.10	0.05	0.21	0.10	0.03	0.13	0.19
plCso	8.50	8.53	8.55	8.46	7.79	8.24	7.59	7.62	8.46	7.57	8.18	7.91	8.24	7.81	7.65	8.04	8.03	7.96	7.65	7.62	7.24	7.36	7.48	7.38	7.35	7.60	7.86	7.48	6.81	8.25	7.21	7.06
Example	25	26	27	28	29	30	31	32	33	34	35	36	37	38	38	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55	56 SFM standard on

:M standard error of the mean of n experiments

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The compounds of Examples 16, 17, 20, 21, 23, 24, 27 and 28 were tested in assay (2) the CD3/VCAM-1 Co-stimulation of T-cell proliferation assay, and the results were obtained as follows:

PIC80	7.4	7.5	6.9	6.9	6.9	7.1	7.5	6.8
Casilpie	16	17	20	21	23	24	27	28

The compounds of Examples 16, 17, 20, 21, 23, 24, 27 and 28 were also tested in assay (3) the Inhibition of lung eosinophil Infiltration and hyper-reactivity in the guinea pig (intratracheal dose given 0.5 hours before and 6 hours after antigen challenge) and the results were as follows:

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Example	Dose	ļul %	% Inhibition
	(µg/kg body	Eosinophil	Hyper-reactivity
	weight)	Accumulation	
16	0.2	62	80
		78	
41	0.2	89	58
	2	61	88
20	0.2		85
	2	79	100
21	0.2	49	82
,	2	79	. 82
23	2	51	79
24	0.2	26.	44
	7	77	85
27	0.2	58	88
	2	08	87
28	0.2	3	70

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•	2	62	47
Dexamethasone	200	55	80
(Positive Control)			

The compounds of Examples 16, 17, 20, 21, 23, 24, 27 and 28 were also tested in assay (4) the RPMI 8866/MAdCAM-1 adhesion assay and the results were as follows:

,u	3	က	2	က	3	က	3	6
SEM*	0.09	0.08	0.16	0.08	0.27	0.05	0.2	0.1
plCso	6.8	6.8	6.7	6.7	7.2	6.6	7.5	6.9
Example	16	17	20	21	23	24	27	28

standard error of the mean of n experiments SEM

Abbreviations

WSCDI

PyBop

1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride

benzotriazol-1-yl-oxy-trispyrrolidinophosphonium hexafluorophosphate

1,3-dilsopropylcarbodilmide DIC

1-hydroxybenzotriazole HOBT

9-fluorenylmethoxycarbonyl tert butoxycarbonyi Fmoc Вос

Cbz

diisopropylethylamine carbobenzyloxy DIPEA

15

dichloromethane DCM

dimethylformamide tetrahydrofuran 보

DMF

1-methyl-2-pyrrolidinone NAP

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understood to imply the inclusion of a stated integer or step or group of integers but not to Throughout the specification and the claims which follow, unless the context requires otherwise, the word 'comprise', and variations such as 'comprises' and 'comprising', will be

the exclusion of any other integer or step or group of integers or steps.

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A compound of formula I:

 \equiv

wherein R' and R2 independently represent

(i) -C, alkyl, -C, cycloalkyl or -C, alkylC, cycloalkyl,

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or such a group in which alkyl or cycloalkyl is substituted by one or more halogen, -CN, nitro, hydroxy or -OC1-ealkyl groups;

(ii) -(CH2),Ar' or -(CH2),DAr';

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or NR¹R² together represent pyrrolidinyt, piperidinyl, piperazinyl, thiomorpholinyl, morpholinyl -CONR¹⁰(CH₂),Ar¹, halogen, -NHSO₂C_{1-s}alkyl, -SO₂NR¹⁰R¹¹, -SO₂C₁₋₈ alkyl or -SO₂Ar² groups; -NR¹º(CO)"(CH₂),Ar¹, -NR¹º(CO)"C₁₃alkylC₃₅ cycloalkyl, -NR¹º(CO)"C₁₅ alkyldiC₃₅ cycloalkyl, or azepinyl, or such a group fused to a benzene ring, optionally substituted by one or more -(CO),(CH₂),Ar', -(CO),C₁₋₈ alkylAr'Ar', -(CO),C₁₋₈alkyl, -(CH₂),OH, -(CH₂),O(CH₂),OH, -(CH₂),OC₁₋₈ alkyl, -O(CH₂),Ar¹, -(CH₂),SO₂Ar¹, piperidin-1-yl, -(CH₂),CONR⁹R⁹, R³ represents -C₁₋₄alkyINHC(=NH)NH₂, -C₂.4alkenyINHC(=NH)NH₂,

or R³ represents -(CH₂),c-2,4-imídazolidinedione, -(CH₂),c(piperidin-4-yl), -(CH₂),c(piperidin-3--(CH₂)₂CHNR¹®CONR®R²¹, -(CH₂)_mNR¹®CONR¹⁴R¹®, -(CH₂)₂NR¹®AP, -(CH₂)₂CONR¹®AP, -C22alkynylNHC(=NH)NH2, -C12alkylNR14R18, -(CH2),CONR14R18, -(CH2),COC12alkyl, -(CH₂),COOR¹¹, -(CH₂),Ar², -O(CH₂),Ar², -(CH₂),CO(CH₂),Ar² or -(CH₂),OAr²;

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yl). -(CH2)。(piperidin-2-yl), -(CH2)。(morpholin-3-yl) or -(CH2)。(morpholin-2-yl) optionally substituted on nitrogen by -(CO),C,alkyl, -(CO),CH2),Ar2 or -C(=NH)NH2; 20

or R³ represents -(CH₂), dibenzofuran optionally substituted by -C₁, alkyl or halogen; or R3 represents -(CH2),-thioxanthen-9-one; R⁴ represents hydrogen, -C₁₋₈ alkyl, -C₁₋₃ alkylC₃₋₈ cycloalkyl, -(CH₂)_qAr², -C₁₋₄alkyl-X-R²,

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R^a represents hydrogen, or R⁴R^a together with the carbon to which they are attached form a -C1-48lkyl SO2C1-4 8lkyl, -C1-68lkylNR12R13 or -C1-4 alkylNR12COC1-6 alkyl; Ce, cycloalkyl ring; Rarepresents hydrogen or -C-alkyl, or Re and R4 together with the N and C atoms to which they are respectively attached form a pyrrolidine ring;

R' represents hydrogen, -(CH2), NR12R13, -(CH2), Ar' or -(CH2), NR12COC14 alkyl; 30

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pyrrolidinyl, piperidinyl, piperazinyl or piperazinyl N-substituted by -C,4 alkyl, -COphenyl or alkyIC34 cycloalkyl, -C24alkenyl or NR8R9 or NR18R17 together represents morpholinyl, Re, Re, Rie and Rindependently represent hydrogen, -Cisalkyl, -Cocycloalkyl, SO₂methyl;

 R^{14} , R^{19} and R^{22} independently represent hydrogen, - C_{16} alkyl, - C_{36} cycloalkyl or -(CH₂), Ar or NR¹⁴R¹⁸ or NR¹⁵R²² together represents morpholinyl, pyrrolldinyl, piperidinyl, piperazinyl or N-R10, R11, R12, R13, R16, R18, R20 and R21 independently represent hydrogen or -C1. alkyl; C1.ealkylpiperazinyl;

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Ar' represents phenyl or a 5 or 6 membered heterocyclic aromatic ring containing 1 to heteroatoms selected from O. N and S optionally substituted by one or more halogen,

Cisalkyi, hydroxy, -OCisalkyi, CFs, nitro, -Art or -OArt groups;

10

Ar² represents phenyl optionally substituted by one or more halogen, -C1.6alkyl, hydroxy, OC, alkyl, -CF, or nitro groups;

Ar represents phenyl, a 5 or 6 membered heterocyclic aromatic ring containing 1 to 3

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-COCH2 CN, -(CH2), NR18R17, -(CH2), NHC(=NH)NH2, -CYNR18(CO), R17, -(CH2), NR15 COR19, heteroatoms selected from O, N or S, or such a group fused to a benzene ring, optionally -(CO)_{*}C₂₋₆ alkenyi, -(CO)_{*}C₂₋₆ alkynyi, -(CO)_{*}C₃₋₆cycloalkyi, -(CO)_{*}C₁₋₆haloalkyi, halogen, substituted by one or more -CO(CH₂), Ard, -(CH₂), Ard, -(CH₂), COArd, -(CO), C₁₋₄ alkyl, -(CH₂),CONR¹⁵R²², -(CH₂),NR¹⁵CONR¹⁵R²², -(CH₂),CONR¹⁶(CH₂),NR¹⁵R²²,

-(CH₂)₆SO₂NR¹⁵R²², -(CH₂)₆SO₂NR¹⁵COAr², -(CH₂)₆NR¹⁵SO₂R¹⁹, -SO₂R¹⁹, -SOR¹⁹, -(CH₂)₂OH -O(CH2),CONR16R17, -O(CH2),COOR15, -O(CH2),OAr2, -O(CH2),Ar2, 3-phenyl-2-pyrazolin-5--COOR15, -CHO, -OC1-10alkyl, -O(CH2),NR16R22, -O(CH2),NHC(=NH)NH2, one or 4,5-dihydro-3(2H)-pyridazinone groups;

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Artrepresents phenyl or a 5 or 6 membered heterocyclic aromatic ring containing 1 to 3 heteroatoms selected from O, N and S optionally substituted by one or more halogen,

25

-C_{1-s}alkyi, hydroxy, -OC_{1-s}alkyl, -CF₃, nitro or -CONH₂ groups;

a, f, k, s and n independently represent 0 or 1;

X and Y independently represent O or S;

b, c, r, x, y and z independently represent an integer 0 to 2; d, g and u independently represent 1 or 2;

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e, h, q and w Independently represent an Integer 1 to 3; and p independently represent an integer 2 to 4; m independently represents an integer 0 to 4;

independently represents an integer 0 to 3;

and salts and solvates thereof.

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2. A compound according to claim 1 wherein R⁴ represents -C₁₋₈ alkyl, R⁵ represents hydrogen or R⁴R⁵, together with the carbon to which they are attached, forms a cyclohexyl ring, and R⁸ represents hydrogen or methyl.

3. A compound according to claim 2 wherein R4 represents -C₁₄ alkyl and R5 and R6 represent hydrogen.

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- 4. A compound according to claim 3 wherein R⁴ represents -CH₂CHMe₂ and R⁵ and R⁴ represent hydrogen.
- 5. A compound according to any one of claims 1 to 4 wherein NR¹R² together represents piperidinyl, piperazinyl, thiomorpholinyl, morpholinyl or 1,2,3,4-tetrahydroisoquinoline optionally substituted by a -(CO)_n (CH₂)_nAr¹, -(CO)_nC₁,3 alkylC₃,6 cycloalkyl, -NR¹0(CO)_nC₁,3 alkylG₃,6 cycloalkyl, -(CH₂)_nOC₁,6 alkyl -(CH₂)_nOC₁,5 alkyli, -(CH₂)_nOC₁,6 alkyli, -(CH₂)_nOC₁,6 alkyli, -(CH₂)_nOC₁,6 alkyli, -(CH₂)_nOC₁,6 alkyli, -(CH₂)_nOC)_nC₁,5 alkylin-1-yl, -(CH₂)_nOC)_nCH₂,5 alkylin-1-yl,5 alkylin-

5

6. A compound according to claim 5 wherein NR¹R² together represents morpholinyl or piperazinyl optionally N-substituted by -(CO)_nC_{1-e} alkyl, piperazinyl N-substituted by -(CO)_n(CH₂)_rAr¹, piperidinyl substituted by -NR¹⁰(CO)_n(CH₂)_rAr¹ or piperidinyl substituted by -(CH₂)_rCONR⁵R⁵.

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7. A compound according to any one of claims 1 to 6 wherein R³ represents -(CH₂)_c-2,4-imidazolidinedione-3-yl, -(CH₂)_c-thioxanthen-9-one-3-yl, -(CH₂)_oAr³, -O(CH₂)_oAr³, -(CH₂)_oOAr³ or -(CH₂)_odibenzofuran.

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- 8. A compound according to claim 7 wherein R³ represents -OCH₂Ar³, -CH₂OAr³ or dibenzofuran.
- 9. A compound according to claim 8 wherein R³ represents -CH2OAr³.
- 10. A compound according to any one of claims 1 to 9 wherein R⁴ and R⁵ have the stereochemical orientation shown in formula (Ia):

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(la)

. A compound of formula (I) which is:

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(2S)-2-{((2S)-2-{[2-(2-Benzoylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[({4-[(2-phenylacetyl)amino}-1-piperidinyl}carbonyl) oxylphenyl}propanoic acid; (2S)-2-({(2S)-4-Methyl-2-{(2-{13-(1-piperidinylcarbonyl)-2-naphthyl}

oxy}acetyl)amino]pentanoyl}amino)-3-{4-[({4-[(2-phenylacetyl)amino]-1-

piperidinyl}carbonyl)oxy]phenyl}propanoic acid;

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(2S)-3-{4-[({4-[(2,2-Dicyclohexylacetyl}amino]-1-piperidinyl}carbonyl) oxy]phenyl}-2-{[(2S)-4-methyl-2-{{2-{4-{(2-{4-{1-piperidinyl}amino}propanoicadid;

(2S)-2-{[(2S)-4-Methyl-2-({2-[4-(1-piperidinylcarbonyl)phenoxy]

2

acetyl}amino)pentanoyljamino}-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl} propanoic acid; (2S)-3-[4-{{[4-{Aminocarbonyl}-1-piperidinyl]carbonyl}oxy)phenyl]-2-{[(2S)-4-methyl-2-([2-[4-(1-piperidinylcarbonyl)phenoxy]acetyl}amino)pentanoyl] amino}propanoic acid; (2S)-3-{4-[({4-[(2-Cyclohexylacetyl)amino}-1-piperidinyl}carbonyl) oxy]phenyl}-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methylpentanoyl) amino]propanoic acid;

(2-10dopnenoxy)acetyljamino}-4-methylpentanoyl) aminojpropanoic acid;
(2S)-3-{4-[((4-[(2,2-Dicyclohexylacetyl)amino]-1-piperidinyl}carbonyl) oxyjphenyl}-2-[((2S)-2-{(2-lodophenoxy)acetyl]amino}-4-methylpentanoyl) aminojpropanoic acid;
(2S)-2-(((2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl}amino)-3-{4-[(4-morpholinylcarbonyl)oxyjphenyl}propanoic acid;

(2S)-2-({(2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl}amino)-3-{4-[({4-[(4-[(2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)oxy] phenyl}propanoic acid; (2S)-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methyl pentanoyl)amino}-3-{4-[(4-[(2-bhenylacetyl)amino}-1-piperidinyl}carbonyl)oxy] phenylacetyl)amino}-1-piperidinyl}carbonyl)oxy] phenylacetyl)amino}-1-piperidinyl}carbonyl)oxy] phenylacetyl

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(2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{((2S)-2-{[2-(2-

iodophenoxy)acetyljamino}-4-methylpentanoyl)aminojpropanoic acid;
(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyljoxy}phenyl)-2-[((2S)-2-{[2-(2-iodophenoxy)acetyljamino}-4-methylpentanoyl)aminojpropanoic acid;
(2S)-3-(4-{[(4-Benzoyl-1-piperazlnyl)carbonyljoxy}phenyl)-2-{((2S)-2-{[2-(2,4-iodophenoxy)acetyl]amino}-1-piperazlnyl)carbonyljoxy

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dichlorophenoxy)acetyljamino}-4-methylpentanoyi)aminojpropanoic acid;
(2S)-3-[4-({[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenylj-2-[((2S)-2-{[2-(2-iodophenoxy)acetyl]amino}-4-methylpentanoyl)aminojpropanoic acid;

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(2S)-2-{[(2S)-2-({2-[2-(Tert-butyl)phenoxy]acetyl}amlno)-4-methyl pentanoyl]amino}-3-[4-({[4-(1-piperidinylcarbonyl}-1-piperidinyl]carbonyl}oxy) phenyl]propanoic acid; (2S)-2-{((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino}-3-[4-({[4-(1-piperidinyl]carbonyl}oxy) phenyl]propanoic acid;

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(2S)-2-({(2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)arnino]-4-methyl pentanoyl}amino)-3-[4-({[4-(1-plperidinylcarbonyl)-1-piperidinyl]carbonyl)oxy) phenyl]propanoic acid;

(2S)-2-{[(2S)-2-({2-[(1-Bromo-2-naphthyl)oxy]acetyl}amino}-4-methylpentanoyl]amino}-3-[4-({[4-(1-piperidinylcarbonyl)-1-plperidinyl]carbonyl} oxy)phenyl]propanoic acid;

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(2S)-2-{[(2S)-2-({2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-(4-{[(4-(2S)-2-[((2S)-2-{[2-(2,4-Dichlorophenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[(4-([(4-fluorobenzyl)amino]carbonyl}-1-piperidinyl) carbonyl]oxy}phenyl)propanoic acid; morpholiny(carbony()oxy]pheny()propanoic acid;

(2S)-2-{((2S)-2-{(2-(2-Benzoylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[(4morpholinylcarbonyl)oxy]phenyl)propanolc acid;

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(2S)-2-{((2S)-4-Methyl-2-{[2-(2-propylphenoxy)acetyl]amino} pentanoyl)amino]-3-{4-[(4morpholinylcarbonyl)oxy]phenyl}propanoic acid; (2S)-2-{[(2S)-2-{(2-{(1-Bromo-2-naphthyl)oxy]acetyl}amino}-4-methylpentanoyl]amino}-3-{4-[(4-morpholinylcarbonyl)oxy]phenyl}propanoic acid;

(2S)-2-{((2S)-2-{[(Benzyloxy)carbonyl]amino}-4-methylpentanoyl) amino]-3-{4-[(4-

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morpholinylcarbonyl)oxyjphenyl)propanoic acid;

(2S)-3-[4-([[4-(2-Furoyl)-1-piperazinyl]carbonyl}oxy)phenyl]-2-[((2S) -2-{[2-(2iodophenoxy)acetyl]amino}-4-methylpentanoyl)amino]propanoic acid;

(2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)aminoj-3-[4-([[4-(2-furoyl)-1-piperazinyl]carbonyl)oxy)phenyl) propanoic acid;

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(2S)-2-{[(2S)-2-((2-[(1-Bromo-2-naphthyl)oxy]acetyl}amíno)-4-methylpentanoyl]amino}-3-[4-(([4-(2-furoyl)-1-plperazinyi]carbonyi]oxy)phenyi] propanoic acid;

(2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy}phenyl)-2-[((2S)-2-{[2-(2-cyclohexylphenoxy)acetyl]amino}-4-methylpentanoyl}amino]propanoic acid;

(2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy)phenyl)-2-[((2S)-(2S)-2-[((2S)-2-{[2-(2-Benzoylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-(4-{[(4-{{2-(4-chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy}phenyl)propanoic acid;

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(2S)-2-{[(2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-(4-{[(4-2-{[2-(2-iodophenoxy)acetyl]amino}-4-methyl pentenoyl)amino]propanoic acid;

(2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidInyl) carbonyl]oxy)phenyl)-2-({(2S)-([2-(4-chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy}phenyl)propanoic acid

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(2S)-3-(4-{[(4-{[2-(4-Chlorophenyl)acetyl]amino}-1-piperidinyl) carbonyl]oxy}phenyl)-2-({(2S)-2-[(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl}amino)propanoic acid;

4-methyl-2-[(2-([3-(1-piperidinylcarbonyl)-2-

naphthyl]oxy}acetyl)amino]pentanoyl}amino)propanoic acid;

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(2S)-2-{[(2S)-2-({2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-{4-[({4-[(2-cyclohexylacetyl)amino]-1-piperidinyl]carbonyl) oxy]phenyl]propanoic acid;

(2S)-2-{[(2S)-2-({2-{2-(Tert-butyl)phenoxy]acetyl}amIno}-4-methyl pentanoyl]amino}-3-{4-[((4-[(2,2-dicyclohexylacetyl)amino]-1-piperidinyl} carbonyl)oxy]phenyl}propanoic acid;

(2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)aminoj-3-{4-{((4-[(2phenylacetyl)amino}-1-piperidinyl}carbonyl) oxyjphenyl}propanoic acid;

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[2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)amlno]-3-{4-[((4-[(2-phenylacetyl)amino]-1-plperidinyl]carbonyl) oxy]phenyl)propanolc acid;

[2S)-3-{4-[({4-[(2-Cyclohexylacetyl)amino]-1-plperidinyl}carbonyl) oxy]phenyl)-2-[((2S)-2-{[2-(2-cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino]propanoic acid; and safts and solvates thereof.

A compound of formula (I) which is:

(2S)-2-[((2S)-2-{[2-(2-lodophenoxy)acetyl]amino}-4-methyl pentanoyl)amino]-3-{4-[(4morpholinylcarbonyl)oxy]phenyl)propanoic acld; (2S)-2-{((2S)-2-((2-[2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-{4-[(4morpholinylcarbonyl)oxy]phenyl}propanoic acid;

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(2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-{[(2S)-2-({2-[2-(tertbutyl)phenoxy]acetyl}amino}-4-methylpentanoyl]amino}propanoic acid; (2S)-2-[((2S)-2-{[2-(2-Cyclohexylphenoxy)acetyl]amino}-4-methyl pentanoyl)amino}-3-{4-[(4morpholinylcarbonyl)oxy]phenyl}propanoic acld;

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(2S)-2-{[(2S)-2-({2-{2-(Tert-butyl)phenoxy]acetyl}amino)-4-methyl pentanoyl]amino}-3-{4-[({4-[(2-phenylacetyl)amino}-1-piperidinyl)carbonyl) oxy] phenyl)propanoic acid;

(2S)-3-(4-{[(4-Benzoyl-1-plperazinyl)carbonyl]oxy}phenyl)-2-{[(2S)-2-({2-{2-(tertbutyl)phenoxy]acetyl}amino)-4-methylpentanoyl]amino}propanoic acid;

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(2S)-3-(4-{[(4-Acetyl-1-plperazinyl)carbonyl]oxy}phenyl)-2-({(2S)-2-{(dibenzo[b,d]furan-4ylcarbonyl)amino]-4-methylpentanoyl}amino)propanoic acid;

(2S)-2-{[(2S)-2-({2-[2-(Tert-butyl)phenoxy]acetyl}amino}-4-methyl pentanoyl]amino}-3-[4-({[4-(2S)-2-(((2S)-2-[(Dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methyl pentanoyl)amino)-3-[4-({[4-(2-furoyl)-1-piperazinyl]carbonyl}oxy)phenyl] propanoic acid;

(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyl]oxy}phenyl)-2-[((2S)-4-methyl-2-{[2-(2-(2-furoyl)-1-plperazinyl]carbonyl]oxy)phenyl] propanolc acid;

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(2S)-3-(4-{[(4-Benzoyl-1-piperazinyl)carbonyljoxy}phenyl)-2-({(2S)-2-[(dibenzo[b,d]furan-4methylphenoxy)acetyl]amino}pentanoyl)amino]propanoic acid; ylcarbony()amino]-4-methyfpentanoy()amino)propanoic acid;

and salts and solvates thereof.

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A compound of formula (I) which is:

(2S)-3-(4-{[(4-Acetyl-1-piperazinyl)carbonyljoxy}phenyl)-2-{((2S)-4-methyt-2-{[2-(2methylphenoxy)acetyl]amino)pentanoyl)amino]propanoic acld;

(2S)-3-[4-([[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-(((2S)-2-

(2S)-3-[4-([[4-(Aminocarbonyl)-1-piperIdinyl]carbonyl}oxy)phenyl]-2-[[(2S)-2-({2-{2-(tert-[(dibenzo[b,d]furan-4-ylcarbonyl)amino]-4-methylpentanoyl}amino) propanoic acid;

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(2S)-2-[((2S)-4-Methyl-2-{[2-(2-methylphenoxy)acetyl]amino} pentanoyl)amino]-3-{4-[(4butyl)phenoxy]acetyl}amino)-4-methylpentanoyi]amino} propanoic acid;

morpholinylcarbonyl)oxy]phenyl)propanoic acid;

(2S)-3-[{[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl}-2-[((2S)-2-{[2-(2benzoylphenoxy)acetyl]amino}-4-methylpentanoyl)amino] propanoic acid; (2S)-2-{[(2S)-2-((2-{4-(Aminocarbonyl)phenoxy]acetyl}amino)-4-methylpentanoyl]amino}-3-[4-([[4-(aminocarbonyl)-1-piperidinyl]carbonyl}oxy) phenyl]propanolc acid;

and salts and solvates thereof.

A compound of formula (I) which is:

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[2S]-3-[4-([[4-(Aminocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-[((2S)-4-methyl-2-{[2-(2methylphenoxy)acetyljamino}pentanoyl)aminoj propanoic acid or a sait or solvate thereof.

A compound of formula (I) according to claim 14 which is:

(2S)-3-[4-([[4-(AmInocarbonyl)-1-piperidinyl]carbonyl}oxy)phenyl]-2-[((2S)-4-methyl-2-[[2-(2methylphenoxy)acetyl]amino}pentanoyl)amino] propanoic acid potassium saft or a solvate

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- A pharmaceutical composition comprising a compound of formula (I) as defined in any one of claims 1 to 15 or a pharmaceutically acceptable salt or solvate thereof in admixture with one or more pharmaceutically acceptable diluents or carriers.
- A pharmaceutical composition comprising a compound of formula (I) according to any one of claims 1 to 15 or a physiologically acceptable salt or solvate thereof in combination together with a long acting eta_2 adrenergic receptor agonist.

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- A compound of formula (I) as defined in any one of claims 1 to 15 or a pharmaceutically acceptable salt or solvate thereof for use as a pharmaceutical.
- Use of a compound of formula (i) as defined in any one of claims 1 to 15 or a pharmaceutically acceptable salt or solvate thereof in the manufacture of a medicament for the treatment of inflammatory diseases.

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A method of treatment or prophylaxis of inflammatory diseases eg. asthma which comprises administering to a patient an effective amount of a compound of formula (I) as defined in any one of claims 1 to 15 or a pharmaceutically acceptable salt or solvate thereof.

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A process for preparation of a compound of formula (I) as defined in any one of claims 1 to 20 which comprises

(a) hydrolysis of a carboxylic acid ester of formula (II)

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wherein R¹, R², R³, R⁴, R⁵ and R⁵ are as defined in claims 1 to 10 and R is a group capable of forming a carboxylic acid ester, or

(b) deprotecting a compound of formula (i) which is protected

A compound of formula (II)

wherein R¹, R², R³, R⁴, R⁵ and Rª are as defined in claims 1 to 10 and R is a group capable of forming a carboxylic acid ester.

A compound of formula (VI)

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wherein P, represents Boc, R4, R6 and R6 are as defined in claims 1 to 4 and 10, and R represents a group capable of forming a carboxylic acid ester.

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A compound of formula (VII)

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wherein P₁ represents Boc, R¹, R², R⁴, R⁵ and R⁰ are as defined in claims 1 to 6 and 10, and R represents a group capable of forming a carboxylic acid ester.

A compound of formula (VIII)

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wherein R¹, R², R⁴, R⁵ and R⁵ are as defined in claims 1 to 6 and 10, HX is a hydrohalic acid and R represents a group capable of forming a carboxylic acid ester.

A compound of formula (XIII)

wherein R⁴, R⁵ and R⁶ are as defined in claims 1 to 4 and 10 and R' represents a hydroxy functionalised polystyrene resin.

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A compound of formula (XIV)

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wherein R3, R4, R5 and R8 are as defined in claims 1 to 4 and 7 to 10 and R' represents a hydroxy functionalised polystyrene resln.

A compound of formula (XXI)

wherein R¹, R², R⁴, R⁵, R⁵ and d are as defined in claims 1 to 6 and 10, R' represents a hydroxy functionalised polystyrene resin and Hal represents halogen,

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INTERNATIONAL SEARCH REPORT

				THE MOUE Application No	
			PCT,	PCT/EP 99/10000	_
A. CLASSIFICATION OF SUBJECT MATE 1PC 7 C070211/58 C C070271/40 C A61K31/445 A	C070295/20 C07K5/06 A61K31/496	C070307/91 C07K17/08 A61K31/5375	C070405/12 C07K1/04 A61P29/00	C07D307/66 A61K31/325	
According to International Patern Classificat	saffication (IPC) or to both	tion (IPC) or to both national classification and IPC	IPC		

Documentation searched other than minimum documentation to the extent that such documents are included in the flaids searched B. FIELDS SEARCHED

Miranum documentation searched (classification system followed by classification symbols)

IPC 7 C070 C07C C07K A61K

Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT

Relevant to claim No. 1, 16, 18–21, 26–28 1,16, 18-21, 26-28 1,16, 18-20 Clation of document, with indication, where appropriate, of the relevant passages WO 98 53817 A (MERK & CO., INC.)
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cited in the application
schemes 1-5
claims 1,11,13; examples WO 98 53814 A (MERCK & CO., INC.)
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scheme 1 WO 98 53818 A (MERCK & CO., INC.)
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cited in the application
claims 1,5,8-10 claims 1.5,154,18-20; examples Category * ⋖ ⋖ Ø

T* later document published after the international (ting date or priority date and not in conflict with the application but clied to understand the principle or theory underlying the invention Patent family members are listed in amex. $[\times]$ Further documents are tsted in the continuation of box C. Special calegories of cited documents; ×

"X" document of particular relevance; the claimed twention cannot be considered novel or cannot be considered to the document is taken alone the document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document defining the general state of the art which is not considered to be of particular relevance
"E" earlier document but published on or after the *International* filling date

"C" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another clistion or other special reason (as specified)

'O' document retenting to an oral disclosure, use, exhibition or other means

document member of the same patent family -To document published prior to the international filing date but later than the priority date defined Date of the actual completion of the International search

Date of mailing of the international search report 29/03/2000 Authorized officer Hass, Name and mailing address of the ISA

European Patent Office, P.B. 5618 Patentiasn 2

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Fax: (+31-70) 340-3016 22 March 2000

page 1 of

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Form PCT/8A/216 (second sheet) (July 1992)

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INTERNATIONAL SEARCH REPORT

Monet Application No

## GB 1 201 121 A (FARBWERKE HOECHST AG) Claims 1,4,5,7; examples		Relevant to claim No.
US 4 105 602 A (R. L. COLESCOTT ET AN B August 1978 (1978-08-08) column 2, 11ne 26 - 11ne 37 columns 3 and 4, bottom of page, coupreaction; columns 5 and 6, top of page coupling reaction W0 98 54207 A (CELLTECH THERAPEUTICS 3 December 1998 (1998-12-03) cited in the application W0 97 03094 A (BIOGEN, INC.) 30 January 1997 (1997-01-30) cited in the application cited in the application		1,16,
W0 98 54207 A (CELLTECH THERAPEUTICS 3 December 1998 (1998–12-03) cited in the application W0 97 03094 A (81005N, INC.) 30 January 1997 (1997–01–30) cited in the application cited in the applicati	•	26-28
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page 2 of 2

INTERNATIONAL SEARCH REPORT

.. remedonal application No.

IIVI EKAYA HONAL SEAKCH KEPORT	PCT/EP 99/10000
Box I Observations where cartain claims were found unsearchable (Continuation of Item 1 of first sheet)	ustion of item 1 of first sheet)
This international Search Report has not been astablished in respect of certain cialms under Article 17(2)(a) for the following reasons:	Article 17(2)(a) for the following reasons:
1. [X] Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claim 20 15 directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged	nemek: he human/animal based on the alleged
2. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically:	he prescribed requirements to such
3. Claims Nos.: because they are dependent dalms and are not drafted in accordance with the second and third sentences of Rule 6.4(s).	id and third sentences of Rule 6.4(s).
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)	2 of first shaet)

	adon, as follows;	
	al Saarching Authority found muttiple inventions in this international application, as follows:	
	This international Searching Autho	

4. Colonal fee, this Authorial fee,	3. As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which faes were paid, specifically claims Nos.:	4. Mo required additional search tees were timely paid by the applicant. Consequently, this internetional Search Report is
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As all required additional search fees were timely paid by the applicant, this International Search Report covers as searchable claims.

The additional search fees were accompanied by the applicant's protest. Remark on Protest

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No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

	ation	2-1998		2-1998	9-1970 0-1970 5-1971			2-1998 3-2000	1 7	∵ .	77	7	7	7	7	7-	~ ~	•	5-1998	∵ !
00001 /66	Publication date	30-1		30-1	30-09 29-10 27-05	50 (5)		30-12 15-0	24-02	10-02	71-05	30-01	16-09	15-04	20-05	05-03	28-09	11-03	25-05-	/n-9'n
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	Patent family member(s)	770319		770329	49666 1593850 1643327	53807		7667498 0984981	716276	6489496	102241	2226868	1193325	9800052	0842196	980033	11511124	98009	324491	3/38
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	Publication date	12-1998	12-1998	12-1998	08-1970		08-1978	12-1998	01-1997			•								1 1 1 1 1 1
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	Patent document clied in search report	9853817	9853814	9853818	1201121	, , , , , , , , , , , , , , , , , , ,	4105602	9854207	9703094											
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